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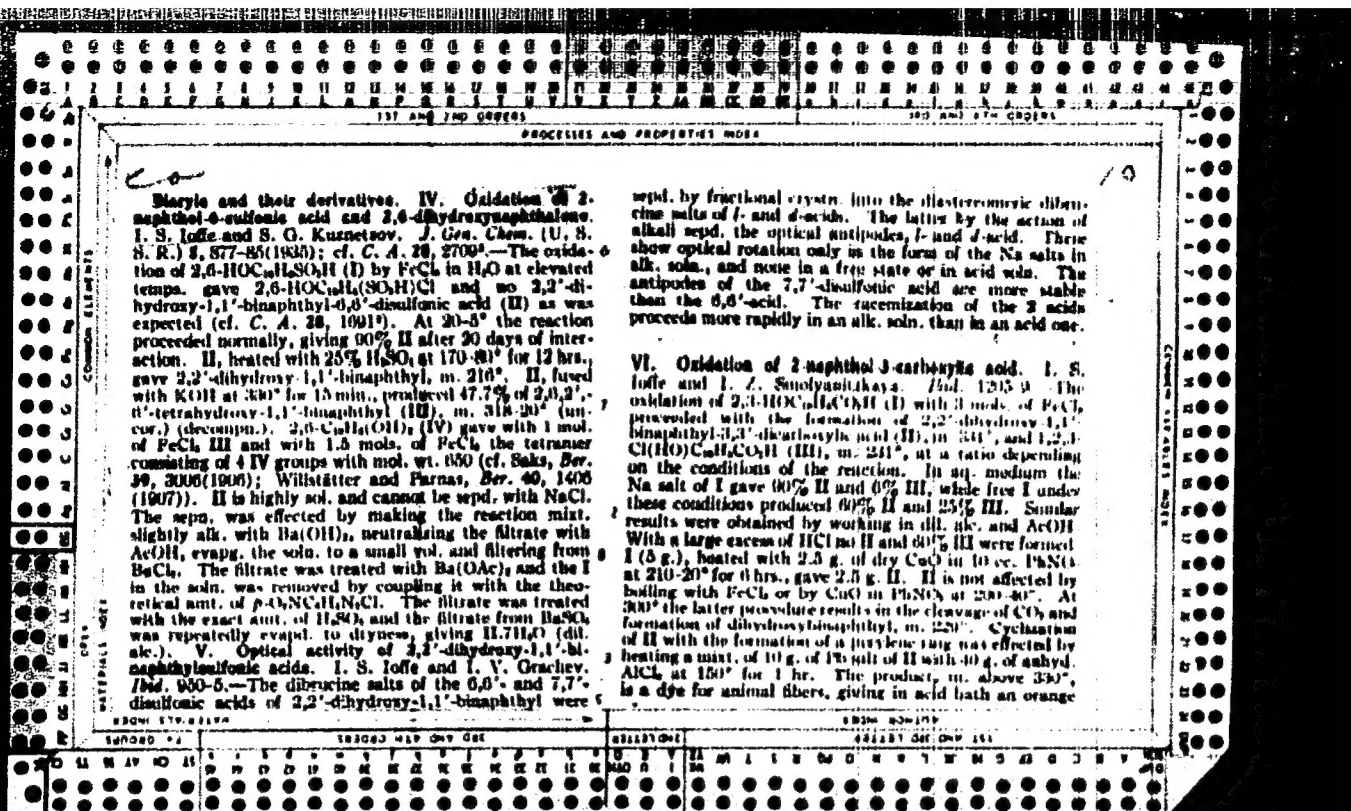
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dyeing changing to a beautiful brown-chocolate, after mordanting with $K_2Cr_2O_7$. The presence of a pyridine ring was demonstrated by the formation of pyridine on distill. with Zn dust. The product is, probably, 1,12-dihydroxyperylene-2-carboxylic acid or 1,12-perylene-2-carboxylic acid. VII. Oxidation of 2-hydroxyanthracene. I. S. 1066. *Ibid.* 1210-12. The oxidation of 2-hydroxyanthracene with $FeCl_3$ in alc. or AcOH resulted in the formation of 2-hydroxy-1,1',0,2'-binaphthyl-ene oxide (I) and a brown compd. of unknown structure. I is unstable, changing at 240° to the brown compd. m. above 300° . I gives with HCl the osmium salt and with Ac_2O in dry pyridine the Ac deriv., m. $247-50^\circ$ (uncr.).
Chas. Blanc

1ST AND 2ND DEGREE		PROCESSING AND PROPERTIES	
<p>CA</p> <p>Summary and their derivatives. VII. Influence of the medium acidity on the interaction of β-naphthol with ferric chloride. I. S. Ioffe, S. G. Kuznetsov and S. Litovskii. <i>J. Gen. Chem.</i> (U.S.S.R.) 3, 1008-9 (1958); cf. C. A. 53, 10489. The effect of the medium acidity on the formation of 2,3'-dihydroxy-1,1'-binaphthyl (I) by the interaction of β-C₁₀H₇OH (II) with FeCl₃ in H₂O (Dionis, <i>J. Russ. Phys.-Chem. Soc.</i> 6, 120 (1874)) was studied by refluxing 0.5 g. II in 1100 cc. of aq. HCl of various concn. with 20 cc. of 1 N FeCl₃ for 2-38 hrs. At a ratio of 20 mols. of HCl to 1 mol. of II the formation of 1,2-C₁₀H₆O₂ (III) begins with 3.4% and gradually rises with the increasing concn. of HCl to 32.4% III at a ratio of 180 mols. of HCl to 1 mol. of II. The formation of I decreases correspondingly from 72.4 to 37.5%. III, m. 70°, was reprecipitated from 1, m. 216° (C₁₀H₆) by steam distillation.</p>			
<p>ASB-3LA METALLURGICAL LITERATURE CLASSIFICATION</p>			
<p>88086 57882194</p>		<p>88086 82-174</p>	
<p>88086 82-174</p>		<p>88086 82-174</p>	

<div style="display: flex; justify-content: space-between;"> 1st AND 2ND COPIES 3RD AND 4TH COPIES </div> <div style="display: flex; justify-content: center; margin-top: 5px;"> PROCESSED AND REPRODUCED </div>									
<div style="position: relative;"> <div style="position: absolute; top: 10px; left: 10px; font-size: 2em; font-weight: bold;">ca</div> <div style="position: absolute; top: 10px; right: 10px; font-size: 2em; font-weight: bold;">10</div> <p style="text-align: center;"> Reaction of <i>p</i>-phenylenediamine and its derivatives with dioxonium salts. I. S. Joffe and V. Ya. Kozlovskikh. <i>J. Chem. Chem.</i> (U. S. S. R.) 6: 977-82 (1936).—In the preliminary experiments, in the presence of substituted metanil yellow for conversion into complex derivatives of <i>p</i>-H₂N·C₆H₄·NHPh (II), the interaction of equiv. units of diphenyl-<i>p</i>-phenylenediamine (III) with N⁺(N₂C₆H₅)₂ (III) resulted in the decoupling of III with a strong evolution of N₂, considerable amt. of unchanged II, a little of diphenyl-<i>p</i>-quinonodiamine (IV) and an am. compd. Evidently II acts as a mild reducing agent and the dioxonium salts as an oxidizing agent: II + RN₂Cl⁺ → IV + RH + HCl + N₂. <i>p</i>-C₆H₄(NH₂)₂ (V) and I react with III similarly with a complete decoupling of III and nearly quant. liberation of N₂. In this decoupling, of derivs. of V, containing at least 1 H atom at each of the NH₂ groups, the formed quinonodiamines react further with the dioxonium salt, forming, probably, arylquinonodiamines: R'N⁺·C₆H₄·NR'' + RN₂Cl⁺ → R'N⁺·C₆H₄·NR'' + (NR'')₂. In the presence of an excess of dioxonium, the decoupling of the latter proceeds further with the formation of probably polyarylsquinonodiamines. <div style="text-align: right;">Chas. Blanc</div> </p> </div>									
<div style="display: flex; justify-content: space-between;"> 1st AND 2ND COPIES 3RD AND 4TH COPIES </div> <div style="display: flex; justify-content: center; margin-top: 5px;"> PROCESSED AND REPRODUCED </div>									

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PROCESS AND PROPERTIES INDEX					100-1000				
BC					100-1000				
<p>New diphenylamine derivatives. J. S. Jorva, S. D. Swenson, and R. G. Swenson (J. Am. Chem. Soc., 1958, 80, 505-507).—2-Chloro-3-nitrobenzoic acid, HCl, H_2O, and HClO_4, in 80% HCl (10 hr., 100-1007) yield the by-products of 4-chloro-3-nitrodiphenylamine-2-sulphonic acid from which the infrared and of the corresponding diacid are reported. This yields a red solid which was coupled with HCl, H_2O, and HClO_4 to give diphenylamine-2-sulphonic acid with boiling H_2O, and 4-chloro-3-nitrodiphenylamine-2-sulphonic acid by the Sandmeyer reaction. R. T.</p>									
<p>ABO-51A METALLURGICAL LITERATURE CLASSIFICATION</p>									
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PROCESSING AND PROPERTIES INDEX

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Stearic and their derivatives. IX. Sulfonation of 2,2'-dihydroxy-1,1'-binaphthyl. I. S. Ioffe, S. G. Kuznetsov and A. A. Panov. *J. Gen. Chem.* (U. S. S. R.) 6, 969-1002 (1936); cf. *C. A.* 30, 3420. --Dissolving 2,2'-dihydroxy-1,1'-binaphthyl (I) with 4 parts of 96% H₂SO₄ on a boiling water bath for 60 hrs. gave 83% 2,2'-dihydroxy-1,1'-binaphthyl-6,6'-disulfonic acid (II), and 13% 2,2'-dihydroxy-1,1'-binaphthyl-6-sulfonic acid (III). II is identical with the acid obtained by oxidation of 2,2'-dihydroxy-1,1'-binaphthyl with FeCl₃ in H₂O (*C. A.* 30, 1047), and HOC₆H₄SO₃H with FeCl₃ in H₂O (*C. A.* 30, 1047). III gives by fusion with KOH 2,2'-dihydroxy-1,1'-binaphthyl (IV), m. 318-20°. The sepn. of II and III depends on the different solubilities of their Ba salts. III fused with KOH at 325° for 15 min. gave 60% 2,2'-dihydroxy-1,1'-binaphthyl, m. 305-7° (sealed capillary). I. S. X. Nitration of 2,2'-dihydroxy-1,1'-binaphthyl. I. S. Ioffe. *Ibid.* 1003-5. --Ten g. I in 200 cc. of 100% AcOH lvs. was treated at 5° with 4.2 g. of 96% HNO₃ in 80 cc. of 100% AcOH, and the mixt. digested on a boiling water bath for 1 hr. The altered ppt. was washed with AcOH and water, then dissolved in 1 N NaOH and reprecip. with HCl, giving 70% 6,6'-dimethoxy-2,2'-dihydroxy-1,1'-binaphthyl (V), m. above 300° (decomp.). V (9.5 g.) in 800 cc. of 1 N NaOH reduced with 180 cc. of 1 N

(1906). --The catalytic reduction of adrenalone (I) to adrenalone (II) was studied by the action of H on 1-HCl in H₂O at room temp. (15-17°) in the presence of Pt, Pd and Ni, with and without carriers. Optimum results of 95% II were obtained at atm. pressure in 24 hrs. with 2% Pd-animal C and in 4 hrs. with 10% Pt-animal C (Mannich, *Arch. Pharm.* 253, 131 (1915)), and in 4 hrs. with Pt black prep. by the method of Loew (*Ber.* 23, 289 (1900)) black prep. by the method of Adams (*Org. Synthesis* 9, 42 (1959)). The reaction with NiCl₂ ignited at 310° was accelerated by the addn. of 1-2 drops of HCl to dissolve some of the pptd. free I, and was completed in 24 hrs. with 70% II. The reduction with 24% Ni-silica gel (heated at 420-500°) at 2 atm. gave 60% II in AcOH and 64% II in alc. in 24 hrs. The ratio of catalysts to I was 1:3 and in alc. in 24 hrs. No improvement of the catalytic action was observed at higher temps., nor in case of Pt and Pd at elevated pressures. Greater activity resulted by increasing the concn. of catalysts in the carriers. Thirty references. Chas. Blum

A S O - S L A METALLURGICAL LITERATURE CLASSIFICATION

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PROCESSING AND PROPERTIES UNIT

REF ID: A60048

CA

Dyeing and their derivatives. III. Synthetic absorption of new dyes derived from dihydroxybenzophenone and tetrahydroxybenzophenone by vegetable fibers. I. S. Isak and M. A. Chigrov. J. Gen. Chem. (U.S.S.R.), 1914-31(1934); cf. C. A. 31, 67N.—Rpts. in the prep. and dyeing of new dyes derived from 2,7,2',7'-tetrahydroxy-1,1'-binaphthyl, 2,6,2',6'-tetrahydroxy-1,1'-binaphthyl, 2,6,2'-trihydroxybenzophenone, and 2,7-and 2,8-dihydroxybenzophenones are described in detail. The monomer dyes were obtained by coupling with p-HO₂C₆H₃NHCl (II), and the dimer dyes with I and p-HO₂C₆H₃NHCl. All 8 dyes gave on wool red dyeings of various shades. Cotton is poorly dyed orange-brown to faint rose suitable to water and soap. Chem. Abstr.

REF ID: A60048

ASB-SLA METALLURGICAL LITERATURE CLASSIFICATION

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1ST AND 2ND ORDERS		PROCESSED AND PROPERTIES INDEX	
<p>Interaction of 1-bromo-2-naphthol with <i>p</i>-nitrobenzoic diacetone oxime. I, S. Ioffe. <i>J. Gen. Chem.</i> (U. S. S. R.) 6, 1074-8 (1936).—In the interaction of 1,2-BrC₁₀H₇ONa (I) with units of <i>p</i>-O₂NC₆H₄N₂OH (II) with displacement of the Br in I by the diacetone component (I + II → O₂N-C₆H₄N₂OH (III) + NaBrO (cf. Hewitt and Mitchell, <i>J. Chem. Soc.</i> 60, 1107 (1908)), no traces of NaBrO could be detected in the filtrate. It is completely decomposed in the process of formation, oxidizing and brominating I, II and III. The resulting para red, m. 140-80°, contains 20.3% Br and is of brown shade. In the presence of Na₂S₂O₄ the destructive action of NaBrO is overcome, giving 80% of bright red III, m. 280-1°. Chas. Blanc</p>		<p>62-1-22</p>	
		<p>ASAC-51A METALLURGICAL LITERATURE CLASSIFICATION</p>	
<p>RECORD SYMBOLS</p>		<p>RECORD SYMBOLS</p>	
<p>RECORD SYMBOLS</p>		<p>RECORD SYMBOLS</p>	

Bromination of 2,7-dihydroxynaphthalene. 1-Bromo-
 and N. M. Fedorova, *J. Gen. Chem.* (U. S. S. R.) 6,
 1070-84 (1930), cf. C. A. 25, 1601, and preceding abstr.
 In the direct bromination of 2,7- $C_{10}H_6(OH)_2$ (I), the iden-
 tification of the position of the Br atoms in the mono-
 and poly-Br derivs. is based on the reaction of aromatic
 diazonium salts ($p\text{-ONC}_6H_4N_2Cl$) in aq. NaOH with
 1,2- $Br_2C_{10}H_6(OH)_2$ with the formation of azo compds.
 by displacement of the Br atom in position 1, and with
 isomeric Br derivs. of $\beta\text{-}C_{10}H_6(OH)_2$ with the formation,
 by coupling, of the usual azo compd. contg. Br (cf. Vesely
 and Surina, C. A. 27, 3230). I is substituted by Br first
 in positions 3 and 8, and then considerably more slowly
 in positions 1 and 6. Contrary to Scholl (*Chem. Zentr.* 3,
 720 (1932)) I with Br gives 3,6-dibromo-2,7-dihydroxy-
 naphthalene and not the 1,8-di-Br isomer. I, m. 143-4°
 (72 g.) in 160 cc. of 100% AcOH at 5-7° was treated with
 liquid Br in definite ratios, and the Br products were puri-
 fied in the usual manner. 3,3',7,7'- $Br_2C_{10}H_6(OH)_2$, m. 135°
 3,6,2,7- $Br_2C_{10}H_6(OH)_2$, m. 144-4° ($C_{10}H_6$, 156-9° (100%
 AcOH). 1,3,6,2,7- $Br_2C_{10}H_6(OH)_2$, m. 203°. Tetra-Br
 deriv. of I, m. 283-4°, resulted only with the aid of an
 $AlCl_3$ catalyst; its structure was not detd. C. B.
2-Methyl-1,9-acenaphthene. Alfredo Darsi and An-
 tonio Semproni, *Gazz. chim. ital.* 66, 183-6 (1930).
 A preliminary note. The mechanism of the reaction be-
 tween β -methylanthracene (I) and $(COCl)_2$ in the pres-
 ence of $AlCl_3$ was studied. I (10 g.), $(COCl)_2$ (30 g.)
 and $AlCl_3$ (7.5 g.) in CS_2 (200 cc.) agitated at 0° for 3

hrs., heated at 100° for 1 hr., decomposed with ice and
 HCl , steam-distd., exd. with dil. Na_2CO_3 (to remove
 β -methylanthraceneacetic acid formed as a secondary
 product), the residue exd. with hot dil. aq. $NaHSO_4$
 and the new residue purified, yield 4 g. of methylene-
 anthracene (II), scarlet, m. 258° (decolorn.) oxidized
 with CrO_3 in AcOH, it yields the acid, m. 206° (decolorn.)
 by Meyer, Fleckenstein and Günther (C. A. 2, 4294).
Monosemicarbazone, ochre-color, m. 278-80° (decolorn.).
 The $NaHSO_4$ soln. from the prepn. of II decomposed
 with HCl yields 1.5 g. of an isomer (III) of II, red, m. 201-8°.
 Oxidized with CrO_3 in AcOH, III yields the acid, m. 204-5°;
 decolorized by Butescu (C. A. 7, 1498). II and Zn dust
 (20 parts) distd. in a current of H, and the solidified prod-
 uct purified with CaH_2 and sublimation, yielded a hydro-
 carbon $C_{14}H_{12}$ (IV), golden yellow, m. 122°, sol. in concd.
 H_2SO_4 . Under the same conditions, III yields a hydro-
 carbon $C_{14}H_{12}$ (V), silky light green, m. 99°; its soln.
 in concd. H_2SO_4 are fluorescent green-yellow. The $C_{14}H_{12}$
 solns. of IV and V show intense blue fluorescence. The
 yields of IV and V were 0.935-0.05 g. from 5 g. of II and
 III, resp. Reduction of II and III by the Kishner-Wall
 method ($H_2NNH_2 \cdot H_2O$) gave no definite products. IV
 is probably 2-methyl-1,9-acenaphthene, while the con-
 stitution of V is less certain. The results obtained in the
 reaction of I, $(COCl)_2$ and $AlCl_3$ should be compared with
 those of Butescu (loc. cit.) and Meyer, Fleckenstein and
 Günther (loc. cit.). Ibid 264.—A correction.

C. C. Davis

ASB-11A METALLURGICAL LITERATURE CLASSIFICATION

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2-Hydroxyanthracene-3-carboxylic acid and its *o*-toluide. I. S. Ioffe and R. A. Shostakhamer. *Org. Chem. Ind.* (U. S. N. R. R.) 6, 267-60 (1937). In the prepn. of 2,3-dihydroxyanthraquinonecarboxylic acid (I) by heating hydroxyterephthaloyl-*o*-benzoic acid with 0.3 mol. of 10% H₂SO₄ at 130-91° for 4.5 hrs. (cf. Kharkharov, *C. A.* 30, 7254), the sepn. of I from the 1,2-isomer (II) is best effected by extg. I with boiling water. By this method a yield of 97% of mixed isomers, m. 265°, was obtained, giving 87% I, m. 285° (292° from 70% AcOH), and 9.1% II, m. 254-5°. Crude I in 25% NH₄OH reduced with Zn dust (cf. Ger. pat. 557,340 (*C. A.* 27, 311), 604,360 (*C. A.* 29, 815)) yielded 69% 2-hydroxyanthracene-3-carboxylic acid, m. 264° (280° from CCl₄/Cl₂). The crude acid (0.13 g.) in 15 cc. toluene refluxed with 0.08 cc. *o*-toluidine for 10-15 min. and then with the addn. of 0.08 cc. PCl₅ for 5 hrs. gave the *o*-toluide, m. 270° (cf. U. S. pat. 1,960,375, *C. A.* 29, 4907). The product is identical with naphthol AS-GR (IG).

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2-Hydroxyanthracene-3-carboxylic acid and its *o*-toluide. I. S. Ioffe and R. A. Shostakhamer. *Org. Chem. Ind.* (U. S. N. R. R.) 6, 267-60 (1937). In the prepn. of 2,3-dihydroxyanthraquinonecarboxylic acid (I) by heating hydroxyterephthaloyl-*o*-benzoic acid with 0.3 mol. of 10% H₂SO₄ at 130-91° for 4.5 hrs. (cf. Kharkharov, *C. A.* 30, 7254), the sepn. of I from the 1,2-isomer (II) is best effected by extg. I with boiling water. By this method a yield of 97% of mixed isomers, m. 265°, was obtained, giving 87% I, m. 285° (292° from 70% AcOH), and 9.1% II, m. 254-5°. Crude I in 25% NH₄OH reduced with Zn dust (cf. Ger. pat. 557,340 (*C. A.* 27, 311), 604,360 (*C. A.* 29, 815)) yielded 69% 2-hydroxyanthracene-3-carboxylic acid, m. 264° (280° from CCl₄/Cl₂). The crude acid (0.13 g.) in 15 cc. toluene refluxed with 0.08 cc. *o*-toluidine for 10-15 min. and then with the addn. of 0.08 cc. PCl₅ for 5 hrs. gave the *o*-toluide, m. 270° (cf. U. S. pat. 1,960,375, *C. A.* 29, 4907). The product is identical with naphthol AS-GR (IG).

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2-Hydroxyanthracene-3-carboxylic acid and its *o*-toluide. I. S. Ioffe and R. A. Shostakhamer. *Org. Chem. Ind.* (U. S. N. R. R.) 6, 267-60 (1937). In the prepn. of 2,3-dihydroxyanthraquinonecarboxylic acid (I) by heating hydroxyterephthaloyl-*o*-benzoic acid with 0.3 mol. of 10% H₂SO₄ at 130-91° for 4.5 hrs. (cf. Kharkharov, *C. A.* 30, 7254), the sepn. of I from the 1,2-isomer (II) is best effected by extg. I with boiling water. By this method a yield of 97% of mixed isomers, m. 265°, was obtained, giving 87% I, m. 285° (292° from 70% AcOH), and 9.1% II, m. 254-5°. Crude I in 25% NH₄OH reduced with Zn dust (cf. Ger. pat. 557,340 (*C. A.* 27, 311), 604,360 (*C. A.* 29, 815)) yielded 69% 2-hydroxyanthracene-3-carboxylic acid, m. 264° (280° from CCl₄/Cl₂). The crude acid (0.13 g.) in 15 cc. toluene refluxed with 0.08 cc. *o*-toluidine for 10-15 min. and then with the addn. of 0.08 cc. PCl₅ for 5 hrs. gave the *o*-toluide, m. 270° (cf. U. S. pat. 1,960,375, *C. A.* 29, 4907). The product is identical with naphthol AS-GR (IG).

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2-Hydroxyanthracene-3-carboxylic acid and its *o*-toluide. I. S. Ioffe and R. A. Shostakhamer. *Org. Chem. Ind.* (U. S. N. R. R.) 6, 267-60 (1937). In the prepn. of 2,3-dihydroxyanthraquinonecarboxylic acid (I) by heating hydroxyterephthaloyl-*o*-benzoic acid with 0.3 mol. of 10% H₂SO₄ at 130-91° for 4.5 hrs. (cf. Kharkharov, *C. A.* 30, 7254), the sepn. of I from the 1,2-isomer (II) is best effected by extg. I with boiling water. By this method a yield of 97% of mixed isomers, m. 265°, was obtained, giving 87% I, m. 285° (292° from 70% AcOH), and 9.1% II, m. 254-5°. Crude I in 25% NH₄OH reduced with Zn dust (

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<p><i>ca</i></p> <p><i>25</i></p> <p>Azo dyes and their derivatives. XIII. Azo dyes from 4,4'-diamine - 2,2' - dihydroxy - 1,1' - binaphthyls. <i>S. Inge. J. Gen. Chem. (U. S. S. R.) 7, 1022-3 (1937); cf. C. A. 31, 3291.</i>—When this compd. is diazotized, only 1 diazo group couples. The other is converted to an OH group, even when an excess of the coupling reagent is present. Violet-colored azo dyes are obtained by coupling the amine with 2-naphthol, 2-naphthol-3-carboxylic acid and 2-naphthyl-3,6-dihydroxy acid. H. M. Leksner</p>																																																			
<p>ASD-31A METALLURGICAL LITERATURE CLASSIFICATION</p> <p>SEARCH SYMBOLS</p> <p>INDEXED BY DATE</p> <p>CLASSIFIED</p> <p>REVIEWED</p>																																																			

COMMON ELEMENTS		PERCENTAGES AND PROPERTIES INDEX	
<div style="position: relative;"> OK </div>		<p>Nitryls and their derivatives. XIV. Ring closure in 6,6'-dimino-2,2'-diethoxy-1,1'-binaphthyl. I. S. Loff, and I. S. Gorbun. J. Gen. Chem. (U. S. S. R.) 7, 1102-5(1937); cf. C. A. 31, (1937).—This binaphthyl does not form a perylene deriv. under any condition, probably because of the decrease in reactivity due to the NO₂ groups. If it is treated with strong H₂SO₄ at 40° for 30 min., it gives 6,6'-dimino-1,1'-binaphthyl 2,2'-oxide, sol. only in HClO₄ and pyridine. This compd. can be re- duced to an amine which can be diazotized II. M. L.</p>	
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<p>Reaction of <i>p</i>-phenylenediamine and its derivatives with diazonium salts. II. Reaction of diphenyl-<i>p</i>-phenylenediamine with diazonized metacetic acid and <i>p</i>-chloraniline. I. S. Joffe and E. T. Lemartovich. <i>J. Gen. Chem.</i> (U.S.S.R.) 7, 1113-18(1937); cf. C. A. 31, 6190. — Further study of the interaction of diphenyl-<i>p</i>-phenylenediamine (I) with <i>m</i>-ClH₃C₆H₄SO₃H (II), resulting in the formation of II with liberation of N₂ and no azo compd., showed that I is capable of reacting with a max. of 3 II units. The reaction proceeds in 2 successive stages with the formation of <i>N,N'</i>-diphenyl-1,4-quinonediimine and its 2-mono- and 2,5-bis(<i>m</i>-sulphophenyl) derivs. The inconclusive results of the interaction of I with <i>p</i>-ClC₆H₄N₂Cl require further investigation. Chas. Blum</p>																																																			
<p>ASB-15A METALLURGICAL LITERATURE CLASSIFICATION</p>																																																			
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Diaryls and their derivatives. XV. The reaction of 2-naphthol-3,6-disulfonic acid with salts of trivalent iron. I. B. Ioffe and H. Chernysheva. *J. Gen. Chem.* (U.S.S.R.) 9, 2401 (1937), cf. 1, 31, 7410. When the Na salt of this acid is heated for 18 hrs. with $FeCl_3$, it gives the Na salts of 1-chloro-2-naphthol-3,6-disulfonic acid (I) and 2,2'-dihydroxy-1,1'-binaphthyl-3,6,3',6'-tetrasulfonic acid (II). If the reaction is carried out in the presence of a large excess of HCl , I is almost the only product, and if $Na(OAc)$ is present to remove all the HCl formed, II is the chief product. If the oxidation is carried out with $Fe(SO_4)_3$, only II is formed. The Cl in I is easily split off by $AgNO_3$ or diazonium salts. XVI. The reaction of salts of trivalent iron with 2-naphthol-3-sulfonic acid and 2-naphthol-5,7-disulfonic acid. I. B. Ioffe and V. I. Kobryakova. *Ibid.* 2182-40. When 2-naphthol-5-sulfonic acid is oxidized by $FeCl_3$, even in the presence of excess HCl , or by $Fe(SO_4)_3$, the only product is 2,2'-dihydroxy-1,1'-binaphthyl-5,5'-disulfonic acid. With $FeCl_3$, 2-naphthol-5,7-disulfonic acid gives chiefly 2,2'-dihydroxy-1,1'-binaphthyl-5,7,5',7'-tetrasulfonic acid, but when excess HCl is added, 1-chloro-2-naphthol-5,7-disulfonic acid is also formed. $Fe(SO_4)_3$ gives only the binaphthyl compd. XVII. The reaction of salts of trivalent iron with 2-naphthol-4-sulfonic acid and its derivatives. I. B. Ioffe and M. A. Benidiktova-Pekher. *Ibid.* 2678-80. With $Fe(SO_4)_3$ or $FeCl_3$, even in the presence of excess HCl , 2-naphthol-4-sulfonic acid gives only 2,2'-dihydroxy-1,1'-binaphthyl-4,4'-disulfonic acid. 6-Nitro-2-naphthol-4-sulfonic acid does not react at all with $Fe(SO_4)_3$, and with $FeCl_3$ gives only 6-nitro-1-chloro-2-naphthol-4-sulfonic acid. H. M. Leicester

The reaction of 1-chloro-2-naphthol with *p*-nitrobenzenesulfonamide salts. I, S. Ioffe. *J. Gen. Chem. (U. S. S. R.)*

7, 2087-8 (1937).—While 1-bromo-2-naphthol and (p-NO₂-C₆H₄N)₂SO₂ give entirely para red, 1-chloro-2-naphthol also forms about 40% of a light yellow comp. m. 123° (decamp.). This is either C₁₀H₇ClON:NC₆H₄NO₂ or its quinoid isomer. In the presence of NaOH, a better quality of para red is obtained in this reaction.

H. M. Lohrstei

H. M. Johnston

Biaryls and their derivatives. XVIII. Oxidation of 2-hydroxyanthracene-3-carboxylic acid. I. S. Ioffe and R. A. Shchekhammer. *J. Gen. Chem.* (U. S. S. R.) 7, 2710-11 (1937); cf. C. A. 32, 2112².—When 2-hydroxyanthracene-3-carboxylic acid is heated in HIOAc with at least a 3-fold excess of ferric NH₄ alum, the corresponding biaryl compound is probably formed, but it at once reacts further to give 80% of 3,3'-dihydroxy-1,1'-binaphthyl-2,2'-dicarboxylic acid, which decomposes above 340°. **XIX.** The reaction of 2-hydroxyanthracene with ferric chloride. I. S. Ioffe and I. S. Klym. *Ibid.* 2712-13. The brown product obtained alone with 2-hydroxy-1,1'-binaphthyl-2,2'-dicarboxylic acid, when 2-hydroxyanthracene is oxidized with ferric (C. A. 30, 1048³) is actually the Fe salt of 2,2'-dihydroxy-3,3'-binaphthyl-10,10'-dione. **XX.** A general consideration of the mechanism of the reaction of 2-naphthol and its derivatives with salts of trivalent iron. I. S. Ioffe. *Ibid.* 2715-18.—In these reactions equilibrium exists between a complex of the type $\text{C}_6\text{H}_4\text{OFeCl}_2$, $(\text{C}_6\text{H}_4\text{O})_2\text{FeCl}$, $(\text{C}_6\text{H}_4\text{O})_3\text{Fe}$ and $(\text{C}_6\text{H}_4\text{O})_3\text{Fe}^{3+}$ ($\text{C}_6\text{H}_4\text{O}$). In acid solutions, the formation of the more complex compound is prevented, and Cl can enter the ring to form a Cl-substituted naphthol. In less acid solutions, the complexes tend to form biaryls.

H. M. Leicester

H. M. Lehyer

A 14-514 METALLURGICAL LITERATURE CLASSIFICATION

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Sulfonation (of organic compounds) • I. S. Ioffe. *Org. Chem. Ind. (U. S. S. R.)* 5, 393 (1968). Al. Va. Ilyukovich. *Ibid.* 305 6. V. Ya. Shaposhnikov. *Ibid.* 396. Al. A. Winkler. *Ibid.* 396-7. N. N. Vorobtsov. Jr. *Ibid.* 397-8. I. I. Vorontsov. *Ibid.* 398 9. V. N. Chumtsev. *Ibid.* 399-400. A. I. Kozlov. *Ibid.* 400. N. N. Vorobtsov. *Ibid.* 400 1. A summary of the joint discussion of the theory and practice of sulfonation of org. compds. at the Prague works Chas. Blaw

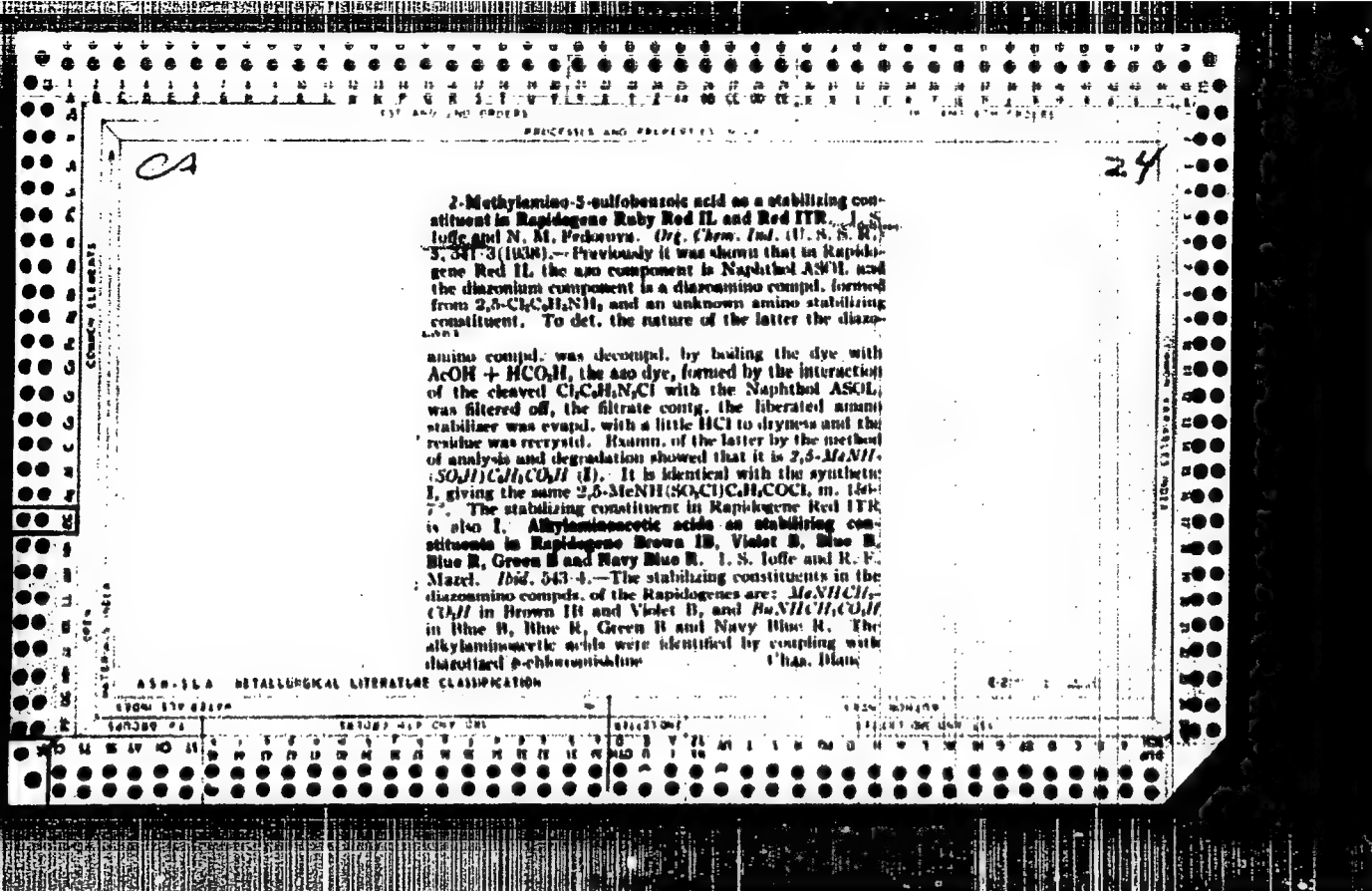
ASB-114 DETAILORICAL LITERATURE CLASSIFICATION

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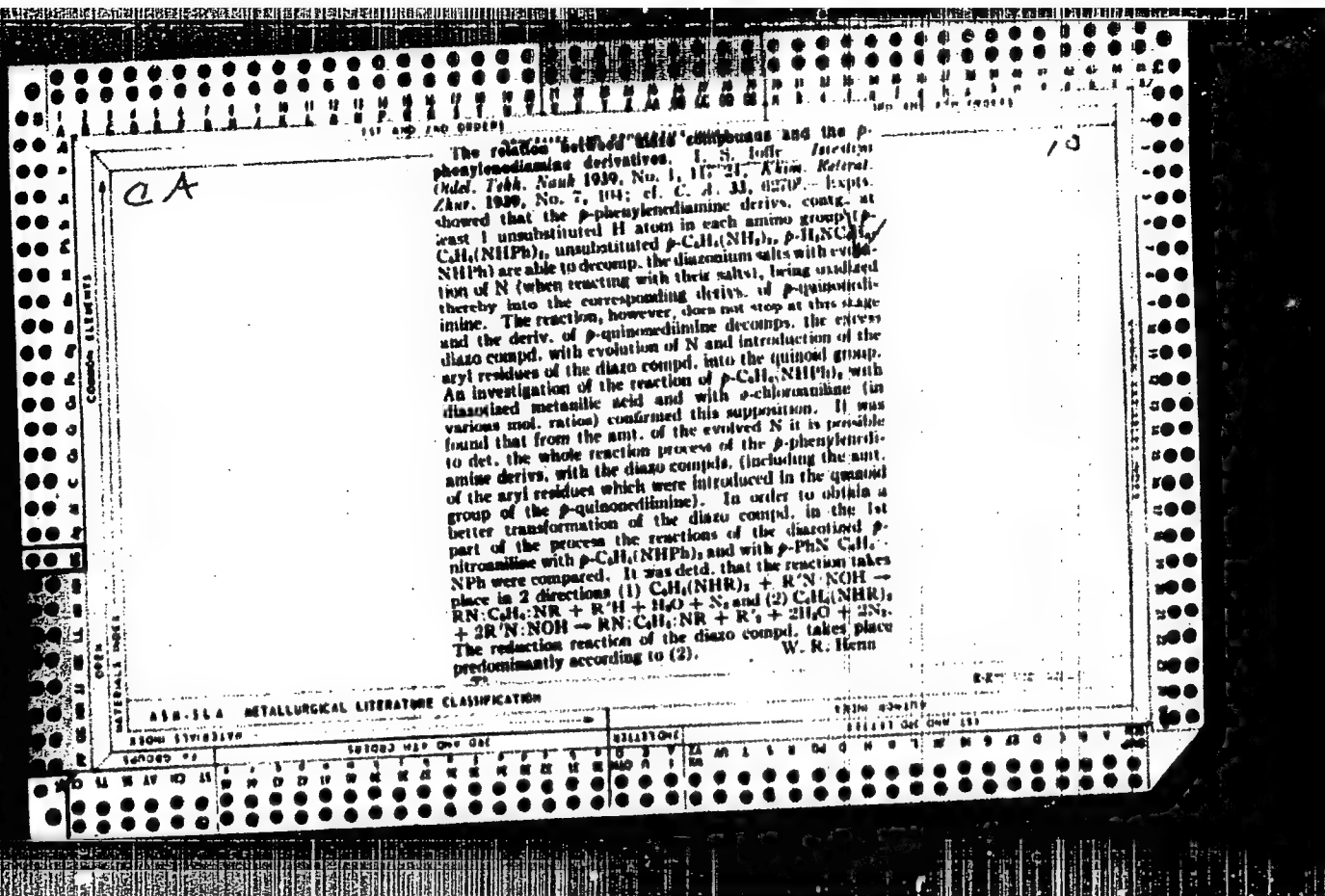
DIASOTIZATION AND PROPERTIES OF DIAZO COMPOUNDS. I. Diazonium compounds and their stable forms. A. I. Poval-Koshits. *Org. Chem. Ind.* (U. S. S. R.) 5, 462-71 (1959).—A summary of the general discussion of the theory and practice of diazotization reactions and the methods of stabilization of diazo compds. at the joint conference of the Tech. Section of the U. S. S. R. Acad. Sci. and the All-Union Mendeleev Chem. Soc. References II. Diazoamine compounds as component parts of the rapidogen dyes. I. S. Ioffe. *Ibid.* 473-6.—A review of American, German, French and Swiss patent literature with approx. 100 references. III. Nitroaminos as starting products for the preparation of ice dyes. M. I. Rozova. *Ibid.* 476-8.—A discussion of the chemistry of nitroaminos and the application of cum. salts of nitroaminos with azo components for rapid dyeing (ice dyeing) based chiefly on literature. IV. Diazo compounds and their use. N. A. Bykhra. *Ibid.* 478-9.—Problems of the production and uses of cum. stabilized nitroamine and diazonium dyes (rapidogen and rapidoids) are discussed. V. The action of mild reducing agents on the diazo compounds. O. M. Golovinski. *Ibid.* 479-84; cf. C. A. 50, 14007.—In the preliminary communication (1) gives theoretical considerations for the proposed expr. table of elec. potentials of various diazonium compds. in neutral, acid and alk. media at various concns. as an aid to volumetric analysis by the oxidation-reduction method. (How Many)

ASM-SLA METALLURGICAL LITERATURE CLASSIFICATION

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<p>Studies in the field of synthesis of diaryl derivatives and their conversion to complex cyclic compounds. I. S. . Inff. Trudy Seriji Akad. Nauk Org. Khim. 1959, 187. 210.—Review of the work done by the author and others in the prepn. of a series of derivs. of 2,2'-dihydroxy-1,1'-binaphthyl from derivs. of 2-naphthol by the action of salts of trivalent Fe. 29 references. B. C. M.</p>																			
<p>ASD-51A METALLURGICAL LITERATURE CLASSIFICATION</p>																			
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PROCESSING AND PROPERTIES INDEX

Sulfonation of 1,2-benzanthraquinone and 1,9-benzanthrone. I. S. Ioffe. *Org. Chem. Ind. (U. S. S. R.)* 6, 95-7 (1959).—Preliminary exper. indicate that 1,2-benzanthrene-7,12-dione (I) and 1,9-benzanthrone (II) react with 100% H₂SO₄ at 20° and with 95% H₂SO₄ at 140-5° with the substitution in the naphthalene nucleus and formation of isomeric monosulfonic acids. I gave 3 acids with the sulfo group in the positions 3, 4 and 5, which were oxidized to 1,2-anthraquinonesulfonic acids. II also gave 3 acids with the sulfo group in the positions 2, 3 and 4, oxidized to anthraquinone-1-carboxylic acid. The sulfonic acids were sepd. as the Ba salts and identified as quinone salts with uncertain res. The results tend to disprove the formation of 1,9-benzanthrone-6-sulfonic acid from II and H₂SO₄, as claimed by Lauer and Iie (C. A. 30, 5967). Ph-1-naphthyl ketone (1-benzoylnaphthalene) reacts with fuming H₂SO₄ at 20° to give 1-benzoylnaphthalene-5-sulfonic acid, identical with the product obtained from the ketone with ClSO₃H in PhNO₂ by Danowski and Moser (C. A. 30, 131). Chav. Blanc

ASAC 114 METALLURGICAL LITERATURE CLASSIFICATION

1. YERSHOV, A.P., IOFFE, I.S.

2. USSR (600)

"The Reaction with Diazo Compounds of Primary Aromatic Amines Containing Salt-Forming Groups -- I. The Tautomerism of Triazenes", Zhur. Obshch. Khim., 9, No. 24, 1939. Sci. Inst. of Organic Intermediate Products and Dyes imeni Voroshilov. Received 7 July 1939.

9. ~~report~~ report U-1626, 11 Jan 1952.

1ST AND 2ND CODES

PRELIMINARY AND PROVISIONAL

10

Reaction of *p*-phenylenediamine and its derivatives with diazonium salts. III. Transformation of diazonium salt. I. S. Ioffe and V. Ya. Solov'chik. *J. Gen. Chem.* (U. S. S. R.) 114 (1939): cf. C. A. 34, 6210. Further study of the interaction of $p\text{-C}_6\text{H}_4\text{N}_2^+\text{Cl}^-$ (I) and its deriva. with diazonium salts (II) showed again that the reaction proceeds in 2 stages in which II, acting as mild oxidizing agents, are decompl. with a nearly complete oxpn. of N, and the formation of the correspond. ing biaryls. In the 2nd stage of the reaction the *p*-quinonediimines, formed by oxidation of I deriva., react further with II to form arylquinonediimines (cf. *ibid.* 114). Thus, the Willstätter imines (cf. C. A. 1, 3001) and their leuco compd. ($\text{PhNHCH}_2\text{NHC}_6\text{H}_4\text{N}_2^+\text{Cl}^-$) react with $p\text{-ONC}_6\text{H}_4\text{N}_2^+\text{Cl}^-$ to yield considerable 1,4'-dinitro-biphenyl, m. 230°, and a product, m. above 330°, probably an aryl deriv. of the imine, which is being investigated. Chas. Blanc

ASTM-51A METALLURGICAL LITERATURE CLASSIFICATION

1ST AND 2ND CATEGORIES		PROCESSING AND PROPERTY INDEX	
<p>New derivatives of diphenyl-p-phenylenediamine. I. S. Joffe and V. Ya. Solovcheik. <i>J. Gen. Chem.</i> (U. S. S. R.) 6, 1448 (1988); cf. preceding abstr. In studies of the structure of intermediates in the formation of aniline black the following new derivatives of diphenyl-p-phenylenediamine were obtained by successive condensation of 1,3,5-trichloro-4-nitrobenzene (I) with PhNHCH₂CH₂NH₂ (II) and PhNH₂, and subsequent reduction and cleavage of the SO₂H group. I was prepd. by sulfonation and nitration of m-C₆H₃Cl₃ (cf. Ger. pat. 120,345). 6-Nitro-3-chlorodiphenyl-p-phenylenediamine-4-sulfonic acid (III) was prepd. in 90.7% yield by refluxing 38 g. of the Na salt of I in 150 ml. H₂O and 20 g. II in 150 ml. alc. with 20 g. Na₂CO₃ for 10 hrs., dilg. with H₂O to 500 ml., filtering and pptg. with 10% HCl. The product dried at 130° resulted in 73% yield by stirring 67 g. III in 200 ml. of 5% Na₂CO₃ into the hot suspension of 55 g. Zn dust, 35 g. NaCl and 6 g. CuSO₄ in 400 ml. H₂O and boiling the mixt. for 2 hrs. After the addn. of a few drops of NaHSO₄, the reaction mixt. is filtered, the filtrate is treated with HCl and the ppt. is dried at a moderate temp. Refluxing 5 g. of the amine with 5 g. ZnCl₂ and 140 ml. of 26% HCl (preliminarily boiled with 1 g. Zn dust) for 3 hrs., dilg. with 100 ml. of cold H₂O and adding an excess of NH₄OH yielded 25% of 6-amino-3-chlorodiphenyl-p-phenylenediamine, m. 148° (benzine). It is a white product, giving colorless soln. in alc., red in C₆H₆, and orange with a green fluorescence in benzine.</p>			
<p>Anticaking 20 g. III (Na salt) in 200 ml. of 50% alc. with 8 g. PhNH₂ and 20 g. Na₂CO₃ at 100-110° for 20 hrs. and acidifying the filtrate with 10% HCl formed 24.0% 6-amino-3-chlorodiphenyl-p-phenylenediamine-4-sulfonic acid, greenish powder, insol. in H₂O. The acid when reduced under the conditions analogous to the preceding expt. yielded 70% of the corresponding amine. The attempts to obtain from II the benzylic Wilitatier leucodiamine by cleaving the SO₂H group produced neg. results. Chas. Blanc</p>			

Sulfonation reaction. IV. Sulfonation of benzan-
threne. I. S. Ioffe and N. N. Mel'teva. *J. Gen. Chem.*
(U. S. S. R.) 9, 1104-6 (1939); cf. C. A. 28, 2384.
Benzanthrene (I) with 30% oleum (87.14% SO₃) is com-
pletely sulfonated in 360 hrs. at 21° and in 6 hrs. at 150°.
Weaker acids (equiv. up to 5% oleum (82.65% SO₃)) are
relatively ineffective at temps. below 150°. I with 22%
oleum for 24 hrs. at 20° gives, after removal of unreacted
I, a mixt. contg. 81% α -benzanthrenesulfonic acid (II) and
19% β -isomer (III), sepd. as the yellow Ba salts by frac-
tional optn. of II from cold water. With I and 30.5%
H₂SO₄ at 150-70° for 8 hrs. the mixt. of sulfonic acids
contains 20% II and 80% III. With 100% H₂SO₄ (IV)
under the same conditions appreciable amts. of disulfonic
acids are obtained, also isolated as the Ba salts. The Ba
salts of both II and III form *quinine salts*, m. 240-2° and
245-2°, resp. Oxidized with Na₂Cr₂O₇ in 30% H₂SO₄ both
II and III give 1-anthraquinonecarboxylic acid, m. 282-4°.
V. Sulfonation of α -naphthyl phenyl ketone. I. S.
Ioffe and G. Z. Naumova. *Ibid.* 1121-3. — α -C₁₀H₇-
COPh (V) readily reacts with 65% H₂SO₄ (VI) at 20°.

At 100° with VI or with oleum at low and high temps. V
is completely converted into water-sol. products. When
heated with VI at 100-70° for 6 hrs. V is hydrolyzed and
sulfonated to give BaOH and CaH₂(SO₃), the latter
isolated from the sulfonation mixt. as the Ba salt. To
prevent hydrolysis V is sulfonated with 10% oleum at 20°
for 24 hrs. to give presumably 1-benzophenanthrene α -
sulfonic acid (VII), whose aniline salt, m. 245°, is identical
with the compd. obtained by Darzenski and Moser
(C. A. 26, 131). With NaOH at 240-320° for 30 min.
VII gives α -naphthol, m. 93°. VI. Sulfonation of 1,2-
benzanthraquinone. I. S. Ioffe and B. N. Kashnitskaya
Ibid. 1124-7. — Contrary to Grachev (*ibid.* 340, 219
(1935)) 1,2-benzanthraquinone (VIII) is more readily sul-
fonated than anthraquinone. VIII with VI at 20° is not
sulfonated after 15 days but at 100° the sulfonation is
practically complete in 6 hrs. IV, as well as weak oleum,
completely sulfonate VIII at 20° in several hrs. VIII
sulfonated at elevated temp. gives a monosulfonic acid
(IX) whose Ba salt, brown, forms a *quinine salt*, brown, m.
118-25°. At low temps. a monosulfonic acid (X) is
obtained whose Ba salt, yellow, gives a yellow *quinine*
salt, m. 202-5°. Oxidation of IX or X with KMnO₄ in
acid soln. gives 1,2-anthraquinonedicarboxylic acid, m.
298°, which indicates that in the sulfonation of VIII the
sulfo group enters exclusively in the side benzo nucleus.

John Livak

ASB-SLA METALLURGICAL LITERATURE CLASSIFICATION

PROCEDURES AND PROPERTIES INDEX

Bromination of 2-hydroxyanthracene. I. S. Joffe,
L. S. Efros and Ts. N. Shcherbova. *J. Gen. Chem.* (U. S.
S. R.) 9, 1129-32(1930).—2-Acetoxyanthracene (I), m.
198°, with Br in cold glacial AcOH gave the 1-Br deriv.
(II), yellow needles, m. 110°, which with NaCr₂O₇ in
hot glacial AcOH gave 2-acetoxyanthraquinone, yellow,
m. 168°, also obtained by oxidation of I. II heated with
dil. NaOH for 10 min. gave 9-bromo-2-hydroxyanthracene,
yellow, m. 112-14°, which with 4-O₂NCH₂NHSO₃ (III)
in alk. soln. gave the 7-oxo deriv., red, m. 254°. 2-Hydroxyanthracene with Br in cold glacial AcOH under-
went both oxidation and bromination to give 7-hydroxy-
1,1'-bianthranyl 9,9'-oxide; 1,10-dibromo-2-hydroxyan-
thracene (IV), yellow, m. 123°, Ac deriv. (V), yellow, m.
198-9°; and 1,9-dibromo-2-hydroxyanthracene, isolated
only as the Ac deriv. (VI), yellow, m. 157-9°. Both V
and VI lose the ac-Br when oxidized with NaCr₂O₇ in
glacial AcOH to give 1-bromo-2-acetoxyanthraquinone, m.
171°. IV with III gave the 1-oxo deriv., red-brown, m.
284°. John Livak

ASB-SLA METALLURGICAL LITERATURE CLASSIFICATION

117 AND 120 OBJECT PROCESSES AND PROPERTIES INDEX

Preparation of nitrophenyl α -naphthyl ketone. I. S. Ioffe and S. S. Bravina. *J. Gen. Chem.* (U. S. S. R.) 9, 1122-3 (1939).—NO₂ derivs. of α -C₁₀H₇CO₂H (II) with the NO₂ group in the Ph nucleus cannot be prepd. by the direct nitration of II. Instead these compounds are prepd. either from the corresponding ONC₆H₄CHO and α -C₁₀H₇CO₂H (C. A. 32, 5037) or from ONC₆H₄COCl (II) and C₁₀H₇ (III) by the Friedel and Crafts synthesis. To m-II (25 g.) with 17.2 g. III in 75 cc. CCl₄ is added 21 g. AlCl₃ at 0° over a period of 2 hrs. to give, after standing for 3 days, 70% of crude α -nitrophenyl α -naphthyl ketone, light yellow (alc.), m. 124°; phenylhydrazones, bright yellow, m. 194°. In the same manner p-II with III gives 68% of the β -nitrophenyl deriv., light yellow, m. 80°; phenylhydrazones, not purified. John Livak

ASS-55A METALLURGICAL LITERATURE CLASSIFICATION

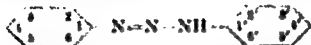
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Macryls and their derivatives. XXI. Oxidation of α -naphthol. I. S. S. Loffe and B. K. Kirichovskiy. *J. Gen. Chem.* (U. S. S. R.), 9, 1130-42 (1939); cf. C. A. 32, 2031f.— α -Naphthol (25 g.) in 2.5 l. boiling water is oxidized with 300-540 cc. 5% FeCl₃ soln., added dropwise and with stirring at 70-80°, to give a mixt. (21 g.) of 4,4'-dihydroxy-1,1'-binaphthyl (I), m. 300°, difficultly sol. in hot benzene, and 1,1'-dihydroxy-2,2'-binaphthyl (II), m. 230°, sol. in hot benzene. I and II with AgCl in pyridine form the *di-Ac* derivs., m. 217° and 196°, resp. Both I and II form azo dyes. I with diazotized *p*-nitroaniline in alk. soln. gives the 3-azo, dark red and the 4,3'-*diaso* deriv., dark red, m. above 330°; II gives the 4-azo, red powder, and the 4,4'-*diaso* compl. (III), brick-red, m. above 330°. Of the 4 dyes only III, having 3 OH groups ortho to the azo groups, is insol. in alkali. I (5 g.) with 28 g. AlCl₃ heated under anhyd. conditions at 150-180° for 3 hrs. undergoes cyclization with the formation of 3,10-perylenequinone, dark brown, m. 350°; and 3,10-dihydroperylene (IV), yellow, m. 227°; di-Bz deriv., m. 265°. IV dissd. with Zn dust gives perylene, orange, m. 260°. II with AlCl₃ is recovered unchanged. II (5 g.) with 20 g. ZnCl₂ under anhyd. conditions at 220-30° for 4 hrs. gives 2.5 g. 2,2'-binaphthyl 1,1'-oxide, grayish yellow, m. 182°, unchanged when dissd. with Zn dust; picrate, m. 173°. I fused with ZnCl₂ under the same conditions remains unchanged.

XXII. Biphenanthryl chloride. I. S. Loffe. *Ibid.* 1148-4.—Oxidation of 2-hydroxyphenanthrene with FeCl₃ in AcOH or EtOH gives 2,2'-dihydroxy-1,1'-biphenanthryl, which, with an equal wt. of AgCl in boiling benzene for 6 hrs., is oxidized to 1,1'-biphenanthryl 2,10',2'-*tri*-oxide (V), bright yellow, m. 290°. V is extremely stable and remains unaltered after treatment with strong oxidizing agents. John E. Livsey

Reaction of diazo compounds with primary aromatic amines containing salt-forming groups. A. Tautomerism of triazines. A. P. Ershov and I. S. Ioffe. *J. Gen. Chem.* (U. S. S. R.) 9, 2211-18 (1939). The following is derived, of diphenyltriazine,



have been prepd.: 3-(7,3'-SO₂Na); 2,5,2'-Cl₂3'-SO₂K;
2,5-Cl₂2'-CO₂Na; 2,5-Cl₂3'-CO₂Na; 2,5-Cl₂4'-CO₂Na;
2,5-Cl₂3'-SO₂Na; 2,3-Cl₂3'-SO₂Na; 2,5-Cl₂4'-SO₂Na;
4-Cl₂-SO₂Na; 2-Cl₂3'-SO₂Na; 4-Me₂-SO₂Na; 4-Me-
4'-SO₂Na; 2,5-Cl₂2',5'-(SO₂Na); 2,5-Cl₂3',5'-(SO₂-
Na); 2,5-Cl₂3'-CO₂H₂4'-SO₂Na; 2,5-Cl₂2'-SO₂H₂4'-
Na; 2,5-Cl₂3'-CO₂H₂3'-SO₂Na; 2,5-Cl₂2'-SO₂H₂3'-

CO₂Na. The prepu. of 2,5-dithio-*d*-sulfodiphenyltriazenes is given as example. A soln. of 0.08 mole 2,5-ChloroC₆H₃N₃NNHPh is added slowly while cooling to a soln. of 0.14 g. Na sulfonate and 10.32 g. NaOAc in 80 ml. H₂O. The reaction product is filtered after 1 hr., the residue washed with H₂O and dissolved in a 5% NaOH soln. The soln. is filtered and the filtrate is acidified with AcOH. The other compds. listed above are prepd. in a similar way. These triazene deriva. exist in 2 tautomeric forms $RN:NHNR' \rightleftharpoons RNHN:NR'$ and the preponderance of one or the other form depends on the character and position of the substituents in the phenyl radicals. III. General course of reaction. *Ibid.* 2210-31.—The following are some deriva. of diphenyltriazene: α -prepd.: 4-Me, 2'-CO₂Na, 4'-SO₃Na; 4-Me, 2'-CO₂Na, 4'-SO₃Na; 4-Me, 2'-SO₃Na; 4-NH₂, 2'-SO₃Na; 4-NH₂, 2'-CO₂H, 4'-SO₃Na. The mechanism of the reaction between diazo compds. and amines, i.e., salt-forming groups was investigated under various conditions. Gertrude Reed

Corrosion Resistance

ASM-55A METALLURGICAL LITERATURE CLASSIFICATION

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PROCESSES AND PROPERTIES INDEX

The characteristic peculiarities of the anthracene derivatives. Investigation of the reaction of 2-hydroxyanthracene with iron chloride. L. B. Ioffe and L. S. Efrem. *Trudy LKKhTI, Lab. Khim. Referat. Zhur.* 1939, No. 6, 68; No. 7, 142-50; *Khim. Referat. Zhur.* 1939, No. 6, 68; cf. C. A. 33, 5904^h.—Two parallel reactions take place with anthracene, the same velocity from the action of FeCl₃ on 2-hydroxyanthracene (I): the oxidation-condensation of 2 mols. of I at the expense of the α-positions with the formation of 2,2'-dihydroxy-1,1'-bianthryl which after formation is immediately further oxidized to 2-hydroxy-1,1'-2'-bianthrylene oxide, and the oxidation-condensation of 2 mols. of I at the expense of their α-positions with the formation of 2,2'-dihydroxy-9,9'-bianthrone. The 1st process is the usual one for α-deriv. of β-naphthol and yields a ruby-colored product which is sol. in benzene, producing a blood-red soln. with a yellow fluorescence (mol. wt. 1064, calcd. 384). The 2nd process, which is specific for anthracene compds., produces a brown product, very little sol. in most org. solvents, sol. in pyridine and insol. in alkali. It is acetylated with difficulty and it dyes wool in a brown color with Cr and Fe mordants.

W. R. Henn

ASH-554 METALLURGICAL LITERATURE CLASSIFICATION

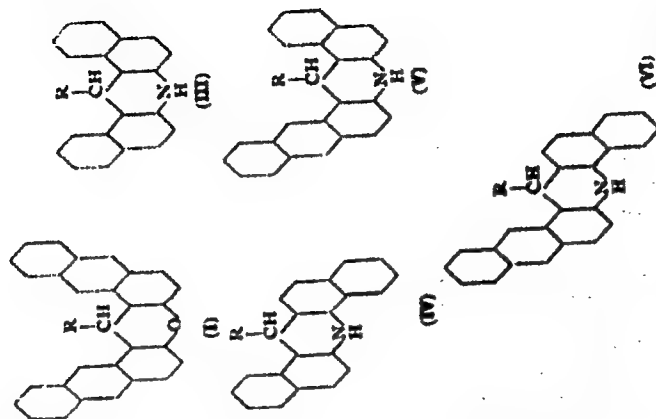
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117 AND 118 ORDERS		PROCEDURES AND PROPERTIES		119 AND 120 ORDERS	
<p>Monosulfonic acids of phenanthrene. (Org. Chem. Ind. (U. S. S. R.) 7, 374-8 (1940).) The following method may be used to sulfonate phenanthrene and to sep. the isomeric monosulfonic acids. Heat 178 g. of phenanthrene gradually to the molten state and then heat to 105-110°. Mix with a stirrer, add from a funnel 150 g. of 92-94% H₂SO₄ in 30 min., regulating the temp. so that it will not rise above 120°, then maintain for 2 hrs. at 110-120° while stirring continuously. Pour into 1 l. of satd. salt soln., wash the vessel and stirrer with the same soln. (the total vol. should be about 1.5 l.), cool and filter. The filtrate is weakly colored, indicating complete absence of disulfonic acids, and is discarded. Wash the light gray ppt. 3 times with satd. salt soln. and compress. The yield is 600-800 g. of paste, depending on the degree of compression. Transfer the paste into 1800 ml. of hot water, heat the mixt. to boiling, boil for 1 hr., cool gradually to 70°, maintain at 70° for 12 hrs., and filter through a heated funnel into a heated receiver, taking care that the temp. does not drop below 60°. The ppt. contains most of the 2-isomer and some unreacted phenanthrene and the filtrate (I) contains most of the 3-sulfonic acid and some 2-isomer. Transfer the ppt. into 2 l. of boiling water, add 5 g. of activated charcoal, boil for 1 hr., filter through a heated funnel, mix, evap. immediately to 300-300 ml., let stand for 12 hrs., filter and dry. The yield was 45-50 g. of pure salt. Boil I, add 10 g. of BeCl₂, boil for 18 min., filter, mix the filtrate and evap. to 1 l. and allow to crystallize in 12 hrs. Filter and dry. The yield was 150-170 g. of sufficiently pure salt. The 2-salts are a mixt. of salts of the 2- and 3-sulfo acids and may be converted to Na salts and sepd. in the usual manner. To convert the sulfo acids into the phenanthranol proceed as follows: Add a few ml. of water to 500 g. NaOH, heat to 280° to melt, add in small amts. 100 g. of 2-sulfonic acid obtained above during the course of 1 hr., raise the temp. to 300° and hold at 300° for 30 min. Pour the melt into 3 l. of water, add while heating HCl to a weakly alk. reaction, filter and add HCl to a weakly acidic reaction. Cool, filter and dry. The product in 100-2° and the yield was 70% of the theoretical. 3-Phenanthranol is obtained under the same conditions except that 350-400 g. NaOH may be used instead of the 500 g. The product usually becomes tarry so that it should be purified by vacuum distn.</p> <p style="text-align: right;">H. Z. Kamich</p>					
<p>ASB-11A METALLURGICAL LITERATURE CLASSIFICATION</p>					
<p>117 AND 118 ORDERS</p>					
<p>119 AND 120 ORDERS</p>					

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157 AND 170 128173		157 AND 170 128173	
<p>2A</p> <p>The interaction of diazo compounds with indophenols.</p> <p>I. S. Jaffe and B. K. Kricheldorf. <i>J. Gen. Chem.</i> (U. S. S. R.) 10:1383-90(1940); cf. <i>C. A.</i> 34, 4730⁹.—An indophenol, obtained by the condensation of $p\text{-NO}_2\text{C}_6\text{H}_4\text{OH}$ with carbazole and having the formula 3-($p\text{-O-C}_6\text{H}_4\text{-N}$)-$\text{C}_{12}\text{H}_9\text{NH}$ (I), was used in glacial AcOH for the reaction with diazotized $p\text{-NH}_2\text{C}_6\text{H}_4\text{SO}_3\text{H}$ (II) at 10-15°. With an excess of II the reaction yielded after 2-3 hrs. 2 isomeric forms of phenylindophenolsulfonic acid, 2 (and 3)-($p\text{-HOS-C}_6\text{H}_4$)-4-O-$\text{C}_6\text{H}_4\text{-NC}_6\text{H}_4\text{-NH}$ (75%), and in the presence of excess of II it yielded after 5-6 hrs. the 2,6-bis-(sulfophenyl) deriv. The monochid was insol. in water but sol. in aq. alk. solns. and the diacid was sol. in water and in aq. alk. solns. The soln. of I in acetone treated with diazotized $p\text{-ClC}_6\text{H}_4\text{NH}_2$ yielded 3 (and 3)-($p\text{-ClC}_6\text{H}_4$)-4-O-$\text{C}_6\text{H}_4\text{-NC}_6\text{H}_4\text{-NH}$ and in the presence of an excess of the reagent the 2,6-bis($p\text{-ClC}_6\text{H}_4$) deriv. Conclusions: The indophenols reacted with the diazo compds., decoupled, them with the liberation of N and adding the aryl residue to the quinoid ring. The arylated indophenols were more colored than the original compds.</p> <p>A. A. Podgorny</p>		10	
<p>ASB-ILA METALLURGICAL LITERATURE CLASSIFICATION</p> <p>FROM STOKESDALE</p> <p>LOGGED BY GUY 101</p> <p>COLLECTOR</p> <p>EXIST ON GUY 101</p>			



in glacial AcOH and H_2SO_4 ; *ms*-(*m*-nitrophenyl)diacetyl-*oxanthrene* (I, $R = m-NO_2C_6H_4$), m. 204° (uncor.); *ms*-(*m*-nitrophenyl)-*oxanthrene* (II), m. 245° ; *ms*-(*m*-nitrophenyl)-*phenanthrene* (III), m. $260-70^\circ$; *p*-nitrophenyl-*phenanthrene* (IV), m. above 330° , was obtained with Br in CH_2Cl_2 yielding a corresponding carbene, m. $319-30^\circ$ (III), with Hg in place of H on the central C atom. The carbene yielded the carbene, m. $264-5^\circ$. Treating the above compound with a mist of $AlCl_3$ and NaCl at 180° yielded the corresponding carbene dyes. The carbene dyes of the same type as for the prep. of II but in which the central C atom was replaced by a nitrogen atom (using *ms*-(*m*-nitrophenyl)-*phenanthrene* as starting material) yielded an isomer of III, *ms*-(*m*-nitrophenyl)-*phenanthrene* (IV), m. 264° . Oxidation of which with Br in $PhCl$ yielded the carbene, m. $264-5^\circ$. On applying the reaction to 2-hydroxyacry-*phenanthrene* (but not in $PhCl$) 2 isomeric *ms*-(*m*-nitrophenyl)-*phenanthrene* derivatives were obtained (V and VI), m. $265-6^\circ$ and 311° . On heating with an $AlCl_3$ -NaCl mist, only V yielded a carbene dye. The properties of the carbene group have stronger base properties than those of the carbene group. A. A. P.

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1ST AND 2ND GROUPS										3RD AND 4TH GROUPS									
PROCESSING AND PROPERTIES INDEX																			
ca										10									
<p>Loffe, I. M.: The Sulfonation of Organic Substances. Leningrad: Naval Med. Acad. 1944. 332 pp. R. 30. Reviewed in Chem. Eng. News 25, 2307(1947).</p>																			
<p>ASAC-SLA. METALLURGICAL LITERATURE CLASSIFICATION</p>																			
<p>SECTION 1</p>										<p>SECTION 2</p>									
<p>SECTION 3</p>										<p>SECTION 4</p>									
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Sulfonation reactions X Sulfonation of benzene

threne and the isolation of its 1'-sulfonic acid. I. S. Joffe and Z. I. Pavlova. *J. Gen. Chem. (U.S.S.R.)* 14, 145-7 (1944) (English summary); *J. C. A.* 49, 927. -When benzanthrene is sulfonated under ordinary conditions, only the 8-sulfonic acid (8) can be isolated. (The numbering used is shown in the structural formula below.)

ASH-SLA METALLURGICAL LITERATURE CLASSIFICATION

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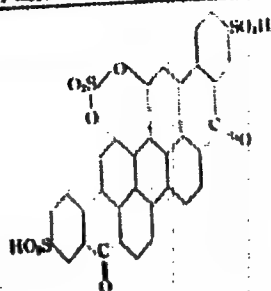
Acetylation of aromatic amines with acetic anhydride in the presence of water. I. S. Ioffe, *J. Gen. Chem. (U.S.S.R.)* 14, 812-15(1944)(English Summary).—It was shown that numerous aromatic amines may be readily acetylated by Ac_2O in aq. medium. PhNH_2 (47 g.) in well condensed by stirring in 140 cc. water and rapidly treated with 10 g. Ac_2O ; the mass warms up and viscosity changes to a thick suspension of AcNHPh , which is filtered on cooling and washed with cold water; yield of acid 80%, m. 114°. Yields of over 80% are obtained under similar conditions from "cellulose water," contg. 31 g. PhNH_2 per l. Yields of better than 80% are obtained with *o*- and *p*- $\text{H}_2\text{NC}_6\text{H}_4\text{Me}$, 1- and 3- $\text{H}_2\text{NC}_6\text{H}_3\text{II}$; MeHNPh , *m*- and *p*- $(\text{H}_2\text{N})_2\text{C}_6\text{H}_3$ give yields of 70-80%. MeHNPh gave a 50% yield. $\text{p-NO}_2\text{C}_6\text{H}_4\text{NH}_2$ gave a low yield of the Ac deriv., probably because of good sol. of the water in water. O. M. Kozolepov

ASB-56A METALLURGICAL LITERATURE CLASSIFICATION
6-477,276,24272

SCHEM SYMBOLISM
FROM SYMBOL

SYMBOL #2
SYMBOL #17 ONV ONE
CELLSTONE

Sulfonation reaction. XI. Sulfonation of violanthrone. I. S. Joffe, F. F. Kekkonen, and L. A. Kalka, J. Gen. Chem. (U.S.S.R.), 18, 810-21(1944) (English summary); *ibid.*, C.A., 39, 2288r, 4567r. —Violanthrone (10 g.) in 100 g. 100% H_2SO_4 was heated to 100° for 8 hrs.; on cooling, the mass was treated with 100 cc. concd. HCl and the ppt. filtered off and purified by reprecip. from water by strong HCl ; 0.5'-violanthronedisulfonic acid so obtained in a deep blue powder, sol. in water and EtOH . The Na salt is readily sol., the Ba salt poorly sol. in water. The above acid (8 g.) in 100 cc. 90% H_2SO_4 was treated with 10 g. MnO_2 at 11° for 4 hrs., after which the mixt. was held at 0° for 24 hrs., poured on ice, and the ppt. dissolved in Na_2CO_3 soln., and treated with hydrazine to reduce the oxazo compd.; the blue soln. was blown with air to oxidize the hemo compd., and the green soln. was acidified with HCl to yield a green ppt. of 2,3'-dihydroxy-0,5'-violanthrone-disulfonic acid. The same compd. was prepd. by sulfonation of 2,2'-dihydroxyviolanthrone by 20% oleum at 20°; it is a green solid, poorly sol. in water and 100% EtOH ; violanthrone (10 g.) was added to 400 g. 20% oleum and heated to 190-210° for 8 hrs.; after treatment with HCl there was obtained the sulfate of 2,3'-dihydroxy-0,5'-violanthronedisulfonic acid.



as a violet powder with benzene extract. The same product is obtained by treatment of 2,4'-dihydroxyxanthone

with 20% above at 100° or similarly from the dihydroxy-
 viomathromedionate acid (see above). Reducing of
 this ester with 10% Na₂CO₃ yields the 2,2'-dihydroxy-
 8,8'-viomathromedionate acid. C. M. Kinsland,
 Crystallization of 8-(p-methoxyphenyl)propionic acid. Wm.
 S. Johnson and Wesley R. Shelberg (Univ. of Wisconsin),
 J. Am. Chem. Soc. 67, 1863-4 (1945). --4-MeOC₆H₄CH₂
 Cl₂CO₂H (I) (3.04 g.) in 100 cc. anhyd. HF in a 14" vessel,
 allowed to stand overnight, gives 94% of unchanged I
 and 3% of 8-methoxy-1-hydrochloride (II); if the reaction
 mixt. is allowed to stand in a closed Cu bomb for 3 days,
 there results 67% of I and 30% of II. The acid chloride
 (prepd. with PCl₅) from 3 g. I and 2.33 g. AlCl₃ in 70 cc.
 CCl₄, stirred 3.5 hrs. at room temp., gave 85% of crude II
 and 4% of crude I; after 0.5 hr. the yield of II was 40%;
 after 20 hrs. it was 60%.

C. J. West

[illegible]

PROCESSING AND PROPERTY INDEX	
CA	<p>Benzenethiopyranic acid. I. S. Ioffe, T. A. Zorina, and V. S. Sushova. <i>J. Gen. Chem. (U.S.S.R.)</i> 14, 955-7(1944) (English summary).—On reduction of 1'-nitrobenzenethrone with hydrochloric acid, there is formed, besides the main product (1'-aminobenzenethrone), a small amt. of 1'-benzenethiopyranic acid, which has indicator properties (yellow changing to blue in strong alkali). The acid is best isolated as follows: (a) alkali filtrate from the aminobenzenethrone is acid. with CO_2, and the yellow acid is added to a small vol. and cooled to yield a yellow ppt. of the above acid, which is purified by crystn. from aq. HCl. The blue (alk.) color of the acid reverts to yellow at pH 11.5. The color changes are explained by suitable tautomeric changes. The acid is poorly stable on heating in solns., esp. in acid media. O. M. Kuznetsov</p>
<p>ASB-11A METALLURGICAL LITERATURE CLASSIFICATION</p>	
<p>EXAM. SYMBOL</p>	
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<p>EXAM. COMMENTS</p>	

1ST AND 2ND SECTIONS										3RD AND 4TH SECTIONS									
PROCESSES AND PROPERTIES INDEX																			
<p><i>ca</i></p> <p>Sulfonation reaction. XIII. Reaction of tertiary aromatic aromatic amines with chlorosulfonic acid. Ioffe and S. S. Bravina. <i>J. Gen. Chem. (U.S.S.R.)</i> 14, 608-73(1944)(English summary); <i>cf. C.A.</i> 39, 2289. Addn. of PhNMe₃ to an equimol. amt. of ClSO₃H yields mainly the anhydro salt, with evolution of HCl and gradual thickening of the mixt.; the product contains only traces of the salt of p-Me₂NC₆H₄SO₃H. Reversal of the addn., however, leads to HCl evolution and thickening of the mixt. only in the initial stage of addn., with the final product constg., in addn. to the anhydro salt, large amts. of the salt of the p-Me₂NC₆H₄SO₃H. The result is explainable by the action of the anhydro salt as a sulfonating agent. O. M. Koudachov.</p>																			
<p>ABN-51A METALLURGICAL LITERATURE CLASSIFICATION</p>																			
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PROCESS AND PROPERTIES INDEX

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Nitro and amine derivatives of dibenzopyrene. I. S. 1086 and L. S. Kirov (1st Leningrad Med. Inst.). *J. Gen. Chem. (U.S.S.R.)* 16, 111-16 (1946).—Dibenzopyrene (6.04 g.) was dissolved with warming in 1200 cc. PhNO_2 and treated, at 25–30°, dropwise with 0.4 g. HNO_3 (d. 1.52) in 10 cc. PhNO_2 ; after 4 hrs. the solvent was concd. to yield 4.9 g. *5-nitrodibenzopyrene*, m. 269–72° (from PhNO_2); boiling with PhNHNH_2 for 15 min. gave *5-aminodibenzopyrene* (I), m. about 310° (from PhCl); *Ac daris*, m. about 350° (from PhNO_2). Dibenzopyrene (3.02 g.) in 100 cc. PhNO_2 treated with 1.4 g. HNO_3 (d. 1.52), gave, in 5 hrs., 3 g. *5,10-dinitrodibenzopyrene*, m. 346–8° (decomp.; from PhNO_2); this, heated for 15 min. with boiling PhNHNH_2 , gave *5,10-diaminodibenzopyrene*, m. 360–8° (from PhNO_2); *di-Ac daris*, m. 440°. Nitration of I in PhNO_2 gave *5-nitro-10-acylamino-dibenzopyrene*, m. 316–20° (from PhNO_2); reduction by boiling with PhNHNH_2 gave *5-amino-10-acylamino-dibenzopyrene*, m. about 357° (from PhNO_2), which on acetylation gave a product identical with that obtained by acetylation of the diamine described above. G. M. Kozolapoff

ASS-314 METALLURGICAL LITERATURE CLASSIFICATION

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<div style="float: left; width: 100px; font-size: 2em; font-weight: bold;">CA</div> <div style="float: right; text-align: right;">10</div> <div style="clear: both;"></div> <div style="text-align: center;"> <p>PROCESSES AND COMPOUNDS INDEX</p> <p>Hydroxy derivatives of triphenylmethane. I. S. Ioffe. U.S.P. 2,683,112, Apr. 30, 1947. HCl gas is bubbled through a mixt. of an aromatic aldehyde 1 and phenol 2 mols. The reaction products are sufficiently pure and can be used directly for the production of the corresponding hydroxyfuchsin dyes. M. Hirsch</p> </div> <div style="text-align: center; border: 1px solid black; padding: 10px; margin-top: 10px;"> <p>Purification of hydroxyfuchsin dyes. I. S. Ioffe. U.S.P. 2,683,113, Oct. 31, 1947. To facilitate the separation of these dyes from their bisulfite soles., the process is carried out in the presence of a satd. soln. of NaCl. Cf. C.A. 43. 3627a. M. Hirsch</p> </div>									
<div style="display: flex; justify-content: space-between;"> <div> <p>ASD-56A METALLOGICAL LITERATURE CLASSIFICATION</p> <p>93000 930 01100</p> </div> <div> <p>93000 930 01100</p> <p>93000 930 01100</p> </div> </div>									

CA
Sulfonation reaction. XIV. Sulfonation of 1-phenyl-3-methyl-5-pyrazolone and preparation and properties of sulfonic acids of phenylmethylpyrazolone. I. B. Ioffe and Z. Ya. Khavun. *J. Gen. Chem. (U.S.S.R.)* 17, 523-7 (1947) (in Russian); cf. *C.A.* 40, 2847. — 1-Phenyl-3-methyl-5-pyrazolone (I) (10 g.) in 50 g. 90% H_2SO_4 was heated 1.5 hrs. at 200-20° until the reaction was complete (a few drops of the mixt. in 3-4 cc. H_2O is treated with several drops of $NaNO_2$ and 2-3 cc. K_2O and shaken; completion of the reaction is shown by a colorless H_2O layer and intensely yellow aq. layer), poured into 100 g. ice, and filtered after standing until ppn. was complete, giving 92% 1-*p*-sulfo-phenyl-3-methyl-5-pyrazolone, difficultly sol. in cold H_2O , sol. in hot H_2O , forms in hot aq. soln. with H_2C_4 the Ba salt, which is sol. in H_2O but can be recrystd. from aq. $EtOH$. The acid or the Ba salt with $NaNO_2$ gives the water-sol. nitroso deriv. $p-O_2NC_6H_4N=N$ gives an orange-yellow azo dye which is fairly sol. in H_2O and in alkalis with a red color. The free acid reduces NH_4Ag_2O and gives with $FeCl_3$ a red color which fades on heating or acidification with HCl . It is stable to hydrolysis; heating to 100-20° with 24% HCl does not affect it. I (10 g.), thoroughly dry, was added slowly to 30 g. 20% oleum and the soln. kept several days at 10-15° (end of reaction shown by treating a few drops with 2-4 g. ice, shaking with Et_2O , and treating the Et_2O layer with $NaNO_2$ and acid, which should give a barely yellow color); after pouring on 100 g. ice, keeping the temp. below 2-3°, the soln. was neutralized with $BaCO_3$, filtered, concd. to 100 cc., again filtered, treated hot

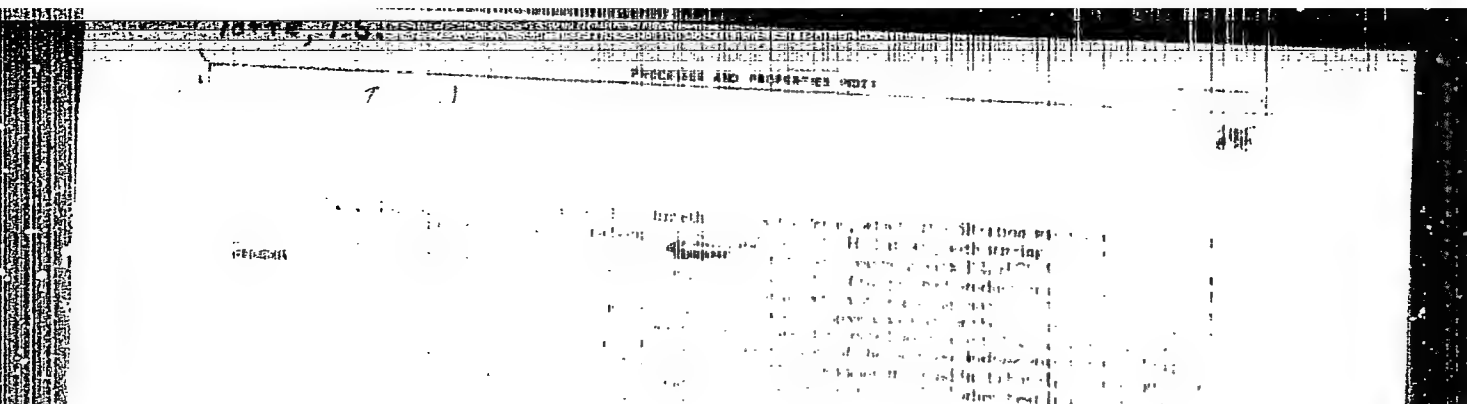
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with $EtOH$ until crystal. began, and cooled, giving 60% 1-phenyl-3-methyl-5-pyrazolone-4-sulfonic acid as white scales; the salt, silvery crystals, contains 5.5 mols H_2O and cannot be completely dehydrated without decomposition. In H_2O , poorly sol. in dil. $EtOH$; $AgNO_3$ gives an insol. Ag salt; $FeCl_3$ gives a blue-violet color; $NaNO_2$ in acid soln. causes loss of H_2SO_4 and gives 4-nitroso-1-phenyl-3-methyl-5-pyrazolone, m. 147-5° (from H_2O); reaction with $p-O_2NC_6H_4N=N$ in the presence of $NaOAc$ (Cals), which is identical with the product of similar coupling of I. The free acid loses the SO_3H group as easily as oleum with cooling, then heated on a steam bath 2 hrs. (test as above should give an almost colorless Et_2O layer), poured slowly with cooling into 200 g. ice, the cooled soln. neutralized with $BaCO_3$, filtered, and the filtrate evapd. to 50 cc. and treated with $EtOH$ to give 78.7% of the Ba salt of 1-*p*-sulfo-phenyl-3-methyl-5-pyrazolone-4-sulfonic acid; on crystal. from dil. $EtOH$ this is obtained as colorless needles, sol. in H_2O , more sol. in dil. $EtOH$ than the Ba completely dehydrated without decomposition and cannot be $AgNO_3$ gives an insol. Ag salt; with $FeCl_3$ it gives a red color; $NaNO_2$ in acid soln. splits off the 4-sulfo group and gives a nitroso compd. identical with that of the *p*-sulfo deriv.; $p-O_2NC_6H_4N=N$ also cleaves a 4-sulfo group and gives an azo dye identical with that from the *p*-sulfo acid. Heating with acids leads to ready loss of the 4-sulfo group. The $NaNO_2$ reaction with the Ba salts of the above acids is quant. and may be used as a basis for their analytical estn.

G. M. Kosolapoff

Sulfonation with H_2SO_4 leads to III through a complex series of steps; the primary products are similar to those produced by oleum; these lead to the formation of IV in later stages of the reaction as well as to accumulation of III through hydrolysis of II by H_2O , III being stable to such hydrolysis. IV then substantially loses the 4-sulfate group by similar hydrolysis and the over-all reaction is the formation of III. 1 (10 g.) added with external cooling -10 to 30, 20% oleum in small portions and allowed to stand 7-10 days at 10-18° and hydrolyzed by pouring slowly into 100 g. externally cooled ice, followed by neutralization with BaCO_3 , gave 66% of the Ba salt (V) of II; the mother liquor after further treatment with HNO_3 gave 17% IV (as the Ba salt (VI)) and a small amt. of V. However, when 10 g. 1 in 50 g. 20% oleum was heated to 100° 1.5-2 hrs., cooled, and the soln. divided into (a) which was rapidly poured into 100 cc. H_2O (without cooling), allowed to cool, and yielded 80% III, and (b) which was carefully poured on 100 g. ice with external cooling; the soln. remained clear and, after neutralization with BaCO_3 , filtration of the BaSO_4 , and soln. of HNO_3

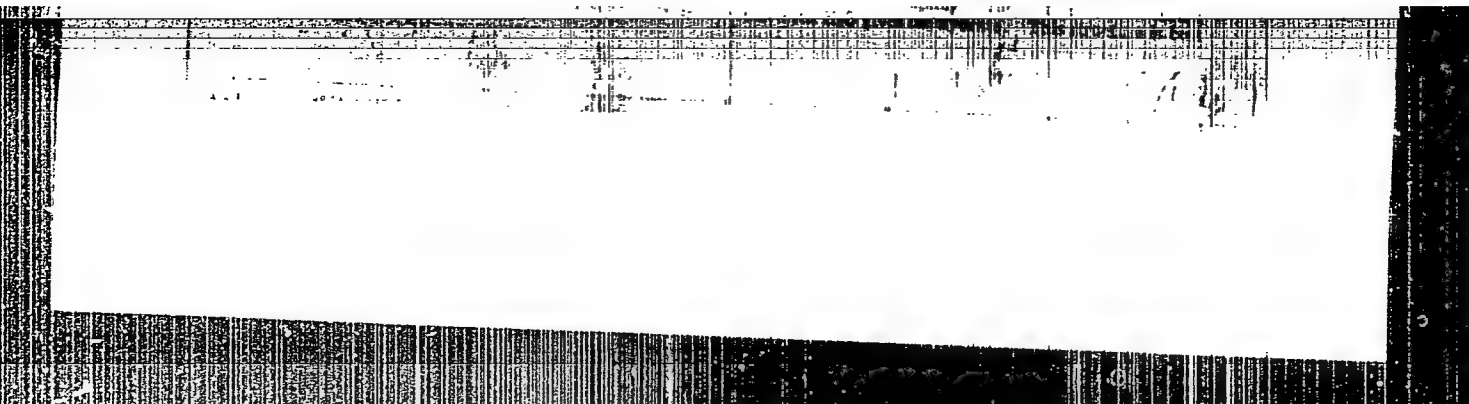
with HCl, giving 61% 5-hydroxy-3-methoxy-2-quinoline (III), m. 230° (decomp.). III, m. 230° (decomp.), is also obtained in 95% yield when 100 g. II (decomp.) is refluxed with 50 cc. HCl in 11. 1/2 hrs. The Ag salt of III refluxed with MeI in MeOH 4 hrs. gives 8.5% II, light yellow needles, m. 123-5°. Slow addn. of 41 g. III to 150 cc. POCl₃ at 80°, refluxing (the mixt.) 1.5 hrs., and pouring into 700 cc. concd. NaOH and ice gives 85% 8-chloro-6-methoxy-3-antiquinoline (IV), m. 203-5°. Heating 6-methoxy-3-antiquinoline (IV), m. 203-5°, with 2 hrs. 4.77 g. IV with Cl₂(OH)CH₂ONa in (C₂H₅O)₂, 7 hrs. and pouring the cooled mist. into 324 cc. H₂O gives 5-(2-hydroxyethyl)-3-antiquinoline (V), bright yellow solid, m. 132-3° (lit. debr. m. 70-1°). V is also obtained in 32% yield by heating 2.4 g. IV in 50 cc. (C₂H₅O)₂ with 0.06 g. 78% KOH in 10 cc. MeOH at 105°. Reduction of 7.1 g. V in EtOH with H₂O, at 40 lb. in 3.5 min. gives 81% 5-(2-hydroxyethyl)-2-antiquinoline-3-endo-quinoline (VI), m. 103-4° (picrate m. 174-0°). An intimate mist. of 0.53 g. VI and 12.9 g. P₂NCl₅(CH₃)₂CH₂Br heated 13 hrs. at 115-25° and 2 hrs. at 125-35° in a 3 atm., giving 2-(8-(6-diethylaminoethyl)plumose)-2-methoxy-3-quinolylglycolal, unstable yellow oil, b.p. 155-60°. Heating 2 g. II with 10 cc. concd. HCl in a sealed tube 6 hrs. at 120° gives 1.4 g. 5,6-dihydroxy-3-antiquinoline. HCl, m. 233-6° (cooked throat, decomp.). Nitration of 17 g. m-ClC₆H₄SO₃Na in 100 cc. C₂H₅O and 9 cc. AcOH gives to -5° with 10 g. HNO₃ (d. 1.42) and 9 cc. AcOH in 60° 7.2-8.0 g. 2-chloro-6-nitroantianiline, m. 117-18°, purified with MeOH in MeOH to 100% 2,8-(Nc)C₆H₃(NH₂)₂ m. 127-8° which is converted into 8-chloro-6-antiquinoline g^o, which is converted into 8-chloro-6-antiquinoline (VII) according to Fournneau, *et al.* (C.A. 24, 5301). Treatment of 2 g. VII with 3 g. MeONa in MeOH, 6 hrs. at 60-80° and overnight at room temp. gives 80% 3-methoxy-3-antiquinoline (VIII), yellow crystals, m. 119-5°. Refluxing 1 g. VIII with 50 cc. MeOH and 3 cc. concd. HCl gives 80% 5-hydroxy-3-antiquinoline (IX), compact white crystals, m. 201° (decomp.). IX is also formed in 10% yield by refluxing VII with KOH in EtOH and 11.2% yield by refluxing VIII with KOH in EtOH and 11.2% yield, almost 100% yield, m. 111-115°. Heating

which gives $\frac{1}{2}$ color in alkali. 2-Amino-3-phenylalanine under these conditions gives 85% 2-amino-3-phenylalanine under these conditions, m. 123-4, which gives no color in alkali. H. M. Littler



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1. The first part of the document is a list of the names of the persons who were present at the meeting.

2. The second part of the document is a list of the names of the persons who were present at the meeting.

3. The third part of the document is a list of the names of the persons who were present at the meeting.

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... product obtained ...
cooled, treated with 0.5 l. Am¹⁴C. and with 7 C)

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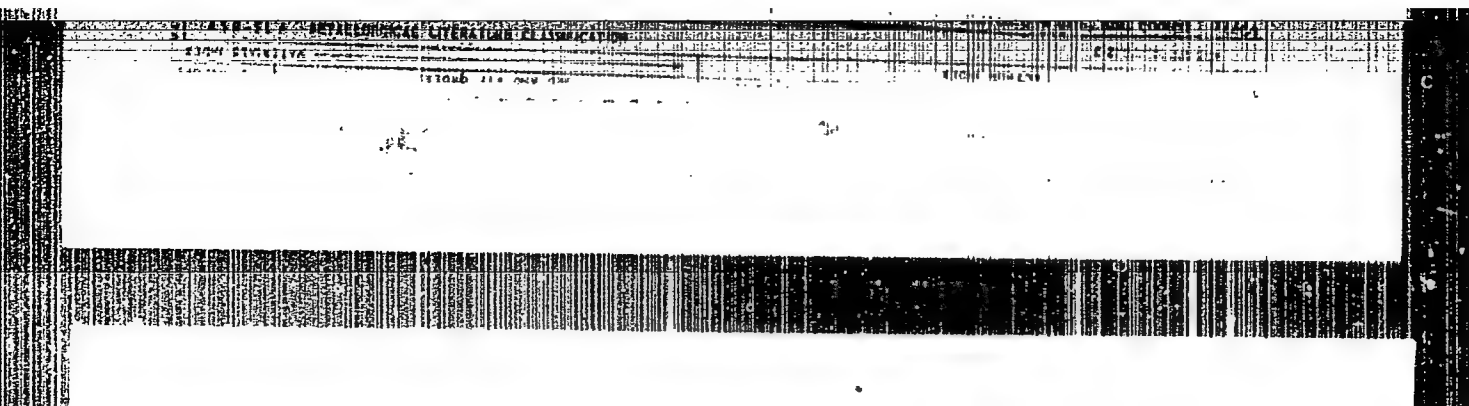
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IOFFE, I. S.

"Investigation in the Field of the Oxyfuchsons Dyes. II. Preparation of 4', 4''-Dioxy-3,3',3(')-Trimetoxyfuchsons (Rubrophens)" (p. 1376)

SO: Journal of General Chemistry, (Zhurnal Obshchei Khimii), 1947, Vol. 17, No. 7

CA
 hydroxyfuchsons dyes. IV. Indicator properties of benzaurin, aurin, and their methoxy derivatives. *Zhur. Khim. (Mikrochim. J.)* (Gen. Chem.) 17, 1001 (1967); cf. C.A. 42, 3434. Titration of solns. of benzaurin (I), dimethoxybenzaurin (II), aurin (III), and trimethoxyaurin (IV) by 0.1 N NaOH potentiometrically shows all 4 substances to be unchanged (pale yellow) up to pH 7; further titration (in which all 4 behave like weak acids), produces a red color going to violet in I, brown-violet in II, pink in III, and violet in IV. I and II show a slight break in the curve, typical of weak acid titrations; III and IV give 2 small breaks indicating 2 phenolic OH groups. The final color stabilization for all 4 substances is reached only at pH 8; on the basis of sharpness of color change, II may have practical uses. However, I and II are not stable at pH about 7, and after 24 hrs. the color formation is much weaker, indicating decomposition; both higher and lower pH values give more stable solns. Further increase of pH beyond 8 results in transformation of the original hydroxyquinone into into colorless quinoid structure, which takes place at pH 11 for I, pH 11.5 for II, in N NaOH for III, and in 1-2 N NaOH for IV. In strongly acid solns. (HCl) these substances also give color changes due to proton addn. and formation of onium-type cations; the changes are: pH 1 for I (orange), pH somewhat under 1 for II (raspberry), pH 2.5 for III (golden) with the best defined color in 5 N HCl, and pH 2 for IV (raspberry). V. Basicity constants of hydroxyfuchsons dyes. *Ibid.* 1010-21. The sodium salts of I-IV were readily prep'd. by the following procedure: a mortar, preheated to 80°, is charged with 50 g. concd. NaHSO₄ soln. in which is gradually

titrated 50 g. of the fuchson derivative. The resulting colorless solid is readily reduced to a powder, which, after soln. in hot H₂O (contg. a drop of 5% AcOH), is filtered and cooled; the products are dried in vacuo over KOH; traces of NH₄ or acid in the atm. must be absent. All of the products are colorless crystals, sol. in H₂O, which develop color on warming in aq. solns. (reversible for brief heating). Acidification of the solns. gives typical ion colors of the anion; soln.: AcOH does not decolor; the basilitic adducts; treatment with alkalis produces colors typical of the alkali salts of the hydroxyfuchson dyes. Analyses indicate equimolar adducts. The decolor. in soln. takes place at pH 4 for acids, and pH 8 for bases for III and IV, and at pH 2 and 7, resp., for I and II. Purification of III may be readily accomplished as follows: crude III (37 g.) at 70° in 600 ml. 1.5% NH₄OH is filtered, the filtrate acidified with AcOH at 70°, and the red ppt. (12.5 g., m. 265°) removed with 12 ml. acid. matter; the filtrate, after H₂O, leaving behind 5 g. insol. matter; with HCl and the soln. of 400 ml. 20% NaCl, is warmed with HCl and the ppt. contg. III HCl treated with NaOAc soln., filtered, and dried, yielding 7.6 g. pure III, m. 202°. Direct conversion of the crude III to the basilitic adduct, followed by HCl treatment, yields III, m. 287-90°. VI. Methoxy- and dimethoxyaurins. I. S. Ioffe and Z. I. Pavlova. *Ibid.* 10, 222-6 (1948). Molten PhOH (40 g.) and 30.4 g. (10, 222-6 (1948)). Molten PhOH (40 g.) and 30.4 g. vanillin were solid. with dry HCl and let stand until crystals started; titration with 2 vols. 30% AcOH gave 30.8 g. 6,6'-trihydroxy-3-methoxyphenylmethane, m. 100-101° (from 30% AcOH), colorless but turning pink in the air. Insol. in petr. ether, sol. in EtOH, AcOH, less sol. in CCl₄. Heating 2 g. with 10 g. AcOH and 4 g. NaOAc 4 hrs. on a

steam bath gave the *tri-Ac deriv.*, colorless needles, m. 112° (from EtOH). *p*-HOC₆H₄CHO (12.5 g.) and 20 g. dissolved in 30 ml. AcOH satd. with dry HCl, allowed to stand several days and the red mass taken up in 10% NaOH, dilut. to 2 l. with water, filtered, and acidified with HCl, gave 30 g. 4,4'-*tri*hydroxy-3,3'-dimethoxytriphenylmethane, m. 175-5.5° (from 30% AcOH), sol. in EtOH, Et₂O, AcOH, AmOAc, less sol. in Cl₂ or petr. ether. *tri-Ac deriv.*, colorless, m. 120-1°. The condensation product of 40 g. PhOH and 20.4 g. vanillin taken up in 150 ml. AmOAc, treated with 40 ml. AmOAc satd. with dry HCl, the soln. treated with 17.5 g. fresh AmONH₂ with stirring over 3-3 hrs., stirred 4 hrs., and let stand overnight gave 70% 3-methoxyvanillin-HCl, purified by heating to 80° with 35 ml. satd. NaHSO₃ soln. and 100 ml. H₂O, pouring into 400 ml. boiling 30% NaCl soln., treating with charcoal, filtering, adding 50 ml. concd. HCl dropwise to the hot filtrate, and collecting the product and washing it with water; the pure product, brownish-tan crystals, decomp. 180°, can be recryst. from AcOH. This HCl salt (12 g.), on addn. to 5 g. NaOAc in water, gives an oil, which on boiling crystallizes and yields hydroxy-*sub*lucosone, m. 274-5°, red crystals with bronze-like tint, can be crystallized from EtOH or AcOH; heating with Ac₂O and NaOAc gives the *tri-Ac deriv.*, yellowish, m. 104-9° (from EtOH), which probably is based on the cardinal structure. 4,4'-*tri*hydroxy-3,3'-dimethoxytriphenylmethane (5 g.) in 25 ml. AmOAc treated with 4 ml. AcOH satd. with HCl, stirred 5 hrs., and let stand overnight gave 3.2 g. 3,3'-dimethoxyvanillin-HCl, which, purified as above, decomp. 165-70°; treatment with NaOAc as above gave the *free dye*, red, m. 231-8° (from EtOH); *tri-Ac deriv.*, colorless, m. 124°. G. M. K.

IOFFE, I. S.

Ioffe, I. S., Pavlova, Z. I., "Research in the Field of Oxyfuchsons Dyes. VI. Methoxy- and Dimethoxyaurins." (p. 222)
(Nav Med Acad imeni Leningard)

SO: Journal of General Chemistry, (Zhurnal Obshchei Khimii), 1948, Volume 18, no. 2

IOFFE, I. S.

22972 Issledovaniye v ob oblasti oksifuksonovykh krasiteley. (Soobshch) ...
9. I. S. Ioffe i A.F. Sukhina. Stroyeniye i tsvetnost' oksimetoksisuksonov.
Zhurnal obshchey khimii, 1949, Vyp. 6, C. 1169-76. Bibliogr: 9 nazv.

SO: LETOPIS' NO. 31, 1949

101 and 102 CROSS
 SUCCESSFUL AND PROPERTIES INDEX
 25
 Hydroxyfuchsons dyes. VII. Monomethoxybenzaurin. I. S. Ioffe and Z. Ya. Khavin. *Ibid.* 1917-28. Molten PhOH (80 g.) spread and chilled over the walls of a long-necked flask was treated with 48 g. Ph₃CCl₂ and the flask was slowly rotated until a uniform mixt. formed; this was kept 15 hrs. and divided into halves; steam distn. of the 1st half, followed by extn. of residue with 5% NaOH and Et₂O, followed by addn. of excess 5% NH₄Cl to the aq. layer gave crude 4-hydroxytriphenylmethane (I), which after drying was rubbed with 6 parts cold EtOH leaving behind insol. 6,6'-dihydroxytriphenylmethane, m. 160-8° (from C₆H₆), coloring violet in air; the di-Ac deriv., m. 101-2° (from dil. EtOH). The product (30 g.) in 150 ml. AmOAc treated with 100 ml. AmOAc satd. with HCl, then with 30 g. AmONO, cooled, let stand overnight, shaken with satd. NaOAc soln. and H₂O, then with 10% NaHSO₃, followed by diln. of the bisulfite ext. by 500 ml. 20% NaCl, filtration with charcoal, and addn. of 200 ml. 15% HCl, gave methoxybenzaurin-HCl, bronze-red crystals, which after washing with NaCl soln., followed by suspension in H₂O and addn. of 30 ml. satd. NaOAc, gave the methoxybenzaurin hydrate, orange-red flakes (70%), m. 60-80°, sol. in EtOH and AcOH with orange color, sol. in alkalis with violet color; evapn. of AcOH soln. of the hydrate in vacuo at 70-80° yields the anhydrous form, red-bronze clusters, m. 158-61° (darkening at 150°) (from xylene). The main product of condensation of 1 mole PhOH, 1 mole PhOH, and 1 mole guaiacol with dry HCl was 6,6'-dihydroxytriphenylmethane, m. 160-1°. VIII. New data on fuchsons and 3-methoxyfuchsons. I. S. Ioffe and Z. Ya. Khavin. *Ibid.* 1917-28. Molten PhOH (80 g.) spread and chilled over the walls of a long-necked flask was treated with 48 g. Ph₃CCl₂ and the flask was slowly rotated until a uniform mixt. formed; this was kept 15 hrs. and divided into halves; steam distn. of the 1st half, followed by extn. of residue with 5% NaOH and Et₂O, followed by addn. of excess 5% NH₄Cl to the aq. layer gave crude 4-hydroxytriphenylmethane (I), which after drying was rubbed with 6 parts cold EtOH leaving behind insol. 6,6'-dihydroxy analog (28%), while addn. of NH₄OH to EtOH soln., followed by H₂O, gave 61% I, m. 158-60° (after leaching with benzene); the 2nd half of the reaction mixt. stirred with 60 ml. satd. warm NaHSO₃ and extd. with C₆H₆ gave a mass of the bisulfite deriv., which, after washing with C₆H₆ and Et₂O, was obtained in 85% yield; this (41 g.) stirred with 100-50 ml. concd. HCl, let stand overnight, and treated with ice gave a red ppt. of *hydrate of the fuchsons*, which on stirring with much H₂O gave 95% yellow I. The dihydroxy deriv. (by-product, above) forms not only in the reaction proper but also during the steam distn. of the mixt. I (25 g.) in 125 ml. AcOH boiled 2 hrs. and cooled, under 90° in vacuo gave on warming with Et₂O 85% yellow fuchsons, m. 100-3° (crude), m. 103-7° (from C₆H₆), Et₂O; an 85% yield results when the bisulfite deriv. is

boiled 4 hrs. with AcOH as above. The bisulfite deriv. is obtained by shaking an alc. soln. of the fuchson with warm satd. NaHSO_3 soln. and diln. to 50% with hot H_2O (I does not undergo this reaction); it is also obtained on addn. of NaHSO_3 soln. to AcOH soln. of fuchson or I; either I or fuchson in excess cold concd. HCl yields the brown HCl salt, which with satd. NaHSO_3 soln. (enough to neutralize residual acid) also gives the same bisulfite deriv. Warm alc. soln. of fuchson is decolorized by a few drops of aq. NH_4OH and on diln. gives I; the same results on boiling with N NaOH and diln. with NH_4Cl soln., or on boiling with 40% AcOH, or on soln. in concd. HCl and diln. Cooled HCl soln. of fuchson-HCl is rapidly hydrolyzed by ice yielding voluminous red fuchson hydrate, m. 59-62°, which on long standing in water yields I; the hydrate is stable only in acid media. Ph_3CCl (192 g.) and 240 g. guaiacol let stand 3 weeks gave a red mass which was divided in halves: the 1st half was ste in distd., powd., and extd. with N NaOH and the soln. treated with CO_2 giving 64% 4-hydroxy-3-methoxytriphenylcarbinol (II), m. 138-9° (from C_6H_6); the 2nd half was stirred on a steam bath with excess satd. NaHSO_3 soln., shaken with warm H_2O and benzene, and the bisulfite deriv. (80%) was ground with concd. HCl and the red soln. of 3-methoxyfuchson-HCl was hydrolyzed by ice after 24 hrs. giving voluminous red hydrate, which on stirring with H_2O gave 78% yellow II, m. 116-18°, pure m. 187-9° (colorless) (from C_6H_6). II (25 g.) boiled 4-5

hrs. with 150 ml. AcOH and evapd. in vacuo gave on extrn. with warm EtOH 7% 3-methoxyfuchson, m. 130-3°, the same was obtained in 30% yield by boiling the bisulfite deriv. with AcOH, evapd., washing with Et_2O , and extrn. with hot C_6H_6 ; the bisulfite deriv. forms under the same conditions as that of fuchson itself and forms a tetrahydrate. The carbinol deriv. of 3-methoxyfuchson: action of NH_4OH or alkali on fuchson give colorless II, m. 157-9°, while boiling 3-methoxyfuchson with 40% AcOH yields yellow carbinol deriv., m. 147°, and hydrolysis of 3-methoxyfuchson-HCl gives yellow hydrate, m. 116-18°, on standing, while the immediate hydrolysis product is a deep-pink solid. Benzhydrol (30 g.), 45 g. PhOH , and 10 ml. AcOH satd. in the cold with HCl and kept several days, steam distd., and treated with CaH_2 gave 4-hydroxytriphenylmethane (73%), m. 108-10° (from dil. AcOH); Ac deriv., m. 82-4°. A similar reaction with 100 g. guaiacol gave 40% 4-hydroxy-3-methoxytriphenylmethane, m. 103-5° (from C_6H_6); Ac deriv., m. 116-18° (from EtOH).

G. M. Kosolapoff

IOFFE, I. S.

PA 67/49T50

Research/Chemistry - Dyes

Fuchsone

May 49

Research in the Field of Oxifuchsone Dyes: VII,
Monomethoxybenzaurin, I. S. Ioffe, Chair of
Org Chem, Nav Med Acad, 6 pp

"Zhur Obshch Khim" Vol XIX, No 5

Synthesized 3-methoxy-4,4'-dioxylphenylmethane
to obtain 3-methoxybenzaurin. It is extracted in
a crystalline form by the passage of dry hydrogen
chloride gas over a saturated mixture containing
proportionally 1 mole of benzaldehyde, 1 mole of
phenol, and 5 moles of guaiacol. The tri-phenyl-
methane derivative, with a melting point of

67/49T50

Research/Chemistry - Dyes (Contd)

May 49

166-1680, remains after the acetyl derivative melts
off at 100-1020. It is then oxidized in an acyl-
acetate solution of amylnitrite to produce the
3-methoxybenzaurin, which has a low melting point
in the hydrate form and melts at 158-160° in the
anhydride form. Alkali solutions of this dye are
violet; strong hydrochloric acid solutions, rasp-
berry; and in the nonionized form, it forms light
yellow solutions similar to the solutions of benza-
urin and its di-methoxy derivative. Submitted
23 Feb 48.

67/49T50

PA 67/49149

IOFFE, IS S.

USSR/Chemistry - Dyes
Fuchsone

May 49

"Research in the field of Oxifuchsone Dyes: VIII,
New Data on Fuchsone and 3-Methoxyfuchsone,"
I. S. Ioffe, Z. Ye. Kozlov, Chair of Org Chem,
Nov Med Acad, 11 3/4 pp

"Zhur Obshch Khim" Vol XIX, No 5

Describes a "new method" for the dehydration of
oxy-triarylcarbinols to produce "good yields" of
fuchsone and 3-methoxy-fuchsone, determining that
1,1-dioxy-tetraarylmethane is also accumulated
in the process. Also describes a "new and easier"

67/49149

USSR/Chemistry - Dyes (Contd) May 49

method for extracting oxy-triaryl-carbinols or
fuchsones from products of the reaction of benzo-
phenonechloride with phenols, and in addition a
"new method" for obtaining oxy-triarylmethanes,
based on the activity of benzhydrol with phenols
in the presence of dry hydrogen chloride gas.
Submitted 23 Feb 48.

67/49149

25

Ca

Hydroxyfuchsons dyes. IX. Structure and color of hydroxymethoxyfuchsons. I. S. Ioffe and A. P. Sukhina. *Zhur. Obshchei Khim.* (J. Gen. Chem.) 19, 1160 (1949); cf. C.A. 43, 8083. Absorption curves are given for fuchsons, 3-methoxyfuchson, benzaurine, mono- and di-3-methoxybenzaurines, aurine, and its mono- and di-3-methoxy derivatives. Fuchsons and their derivatives in unionized state give similar curves in the visible region, but HO and MeO groups cause a small bathochromic shift. Hydroxyfuchsons in mineral acids give curves which are identical regardless of the nature of the substituent. Benzaurines are more intensely colored in acid than alk. solns., than the aurines. MeO groups cause a small bathochromic shift in aurines and benzaurines (in acid or alk. solns.). The most satisfactory alk. soln. was 1% borax, as strong alkalis yield colorless carbinol base anions. G. M. Kosolapoff.

Chair Org. Chem. Naval Med Acad.

ASAC-516 METALLURGICAL LITERATURE CLASSIFICATION

IOFFE, I. S.

Ioffe, I. S. & Khavin, Z. Ya. - "Interaction of methylenic bases of the thiazole series with alkyl halides." (p. 145)

SO: Journal of General Chemistry, (Zhurnal Obshchei Khimii), 1950, Vol. 20, No. 1

CA

Hydroxyfuchson dyes. X. So-called tautomerism of hydroxytriarycarbinols. I. S. Ioffe and Z. Ya. Khavin (Naval Med. Acad., Leningrad). *Zh. Obshch. Khim.* (J. Gen. Chem.) 20, 158-70 (1950); cf. C.A. 43, 9431b. Crit. examn. of exptl. data indicates that Gomberg's school is in error in its defense of "desmotropy" of hydroxytriarycarbinols (C.A. 7, 3307; 10, 82, 33). Heating such compds. as 3-methoxy-4-hydroxytriphenylcarbinol to 75° in porcelain boats in glass app. gave loss of 1.1% only in 50 hrs. from the colored form, while the colorless form lost but 0.6% of its wt.; however, continued heating gave continued wt. loss and formation of sublimable decomposition products; in addn. the "dehydrated" product on treatment with NaHSO_3 soln. gave a ppt. of the initial carbinol, while authentic 3-methoxyfuchson gave a colorless soln. Spectroscopic data of Gomberg and Anderson (C.A. 23, 3457) indicate that the so-called "desmotropes" are probably mixts. of colorless carbinols with fuchsons. Although crystn. of 4-hydroxytriphenylcarbinol and 3-methoxy-4-hydroxytriphenylcarbinol from AcOH of various concns. (40, 60, 80%) gives products of different colors and m.p., quant. colorimetry of their solns. in C_6H_6 merely indicates different amts. of admixture (0.1-1.1%), of corresponding fuchsons to the carbinol, and not individually different substances. This is supported by isolation of similar materials upon crystn. of fuchsons from dil. AcOH of the same concns., i.e. hydration of fuchsons occurring in this case gives mixts. similar to those obtained by dehydration of the carbinols, both reactions being feasible in dil. AcOH . Treatment of such products from 4-hydroxytriphenylcarbinol with dry CaH_2 induces to separate the fuchson impurity; similar sepn. is possible by treatment with NaHSO_3 soln. when the fuchsons dissolve in the form of adducts, while the carbinols are unchanged. G. M. Komlaev

CA

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Hydroxyfuchsones dyes. II. Oxidative alkaline decomposition of hydroxyfuchsones dyes of benzaurine group. I. S. Ioffe. *Zhur. Obshch. Khim.* (1). Gen.

Chem.) 20, 340-55(1944); cf. C.I. 44, 3504. - Acetation of alk. solns. of hydroxyfuchsones dyes gives the corresponding substituted benzophenones. The reaction permitted identification of Dielskott's compl. (*Gost. khim. pat.* 27, 296(1947)), m.p. 131-2° (from EtO) and gualacel) as 4-methoxy-3-hydroxybenzophenone. Only the colored monomeric anions of the dyes are oxidizable; solns. in the form of unoxidized hydroxyquinone deriva. or colorless carbinols are stable to aeration, and the ease of oxidation rises with increased concn. of alkali, although an increase of oxidation-stable carbinol form gives a superficial appearance of lower overall reactivity; low temp. and higher alk. concn. both tend to shift the equil. to the carbinol side. The effect of alkali was confirmed on solns. of 1% Na₂CO₃ and 0.1-5.0% NaOH with aurin, trimethoxyaurin, benzaurin, and dimethoxybenzaurin. Passage of air through 5 g. benzaurin in 250 ml. 1% NaOH on a water bath until red-brown color forms, followed by addn. of 10 ml. AcOH and 2 ml. NaHSO₄ soln. gave 1.5 g. 4-hydroxybenzophenone, m. 134-5°; benzoate, m. 112.5° (from EtOH). Similar reaction with 2,3'-dimethoxybenzaurin gave 3-methoxy-4-hydroxybenzophenone, m. 97-8°. Treatment of this or the 4,5-analog with Me₂SO₄ and 10% NaOH gave 3,4-dimethoxybenzophenone, m. 98-100° (from dil. EtOH). G. M. Kamolajod

Hydroxyfuchsons dyes. XII. Acidic transformations of methoxyfuchsons derivatives. I. S. Ioffe (Naval Med. Acad., Leningrad). *Zhur. Obshch. Khim.* (J. Gen. Chem.), 20, 639-49 (1950); cf. C.A. 44, 6130k.—Fuchsons contg. a MeO group ortho to the CO link undergo an irreversible change on heating with acids; the products

are colorless substances, with the same empirical compo., which are the result of ring closure to *meso*-phenylfluorene derivs.; the distn. of such products with Zn dust should yield *meso*-phenylfluorene, but this has not been accomplished as yet, and the evidence is indirect. Heating 10 g. 3-methoxyfuchson with 100 ml. 15% HCl 1-2 hrs. yields 10 g. brown transformation product, purified by extr. in EtOH and addn. of NaHSO₃ soln., followed by extr. with Et₂O and crystn. from Et₂O. RfOH, 30% AcOH, or C₆H₆. The purified product forms colorless needles, m. 173-4°. poorly sol. in cold aq. alkalies; the product, C₁₈H₁₃O(OH), yields a monoacetate, m. 176°, with Ac₂O-pyridine. 3,3'-Dimethoxybenzaurin similarly heated with 10% HCl yields a colorless product, C₁₈H₁₃O(OH)₂, m. 190-201° (from EtOH and AcOH), which forms a diacetate, m. 227-8° (from AcOH), identical with that formed by heating 3,3'-dimethoxybenzaurin-HCl with Ac₂O and AcCl 2 hrs. on a steam bath. Boiling 3,3'-trimethoxyaurin with 10% HCl 2-3 days gave 5-6 g. transformation product, which was finally purified by boiling in C₆H₆ with Zn dust and AcOH, when it was obtained in 0.5-1.5 g. yield, m. 236-40°; its acetylation product m. 204-5° (from EtOH). The MeO groups in all the aurins investigated are intact in the transformation products, which are believed to contain the OH group in the para position, relative to the central C atom. The Ac deriv. of the last compd. was identical with the acetylation product of rubrocud (cf. Riklin and Postovskii, C.A. 42, 182g), but the structure proposed by R. and P. is questionable.

G. M. Kosolapoff

IOFFE, I. [56.]

[Using engineer Kovalev's method in oil well drilling and operation] Opyt primeneniia metoda inzhenera Kovaleva v burenii i dobyche nefi. 1951.58 p. (MIRA 8:8)

1. Moscow. Tsentral'nyy nauchno-issledovatel'skiy institut mekhanizatsiy truda v neftyanoy promyshlennosti. Byuro tekhniko-ekonomicheskoy informatsii.

(Oil well drilling)

Hydroxyfuchsones dyes. XIII. Acetylation of hydroxy-methoxyfuchsones. I. B. Joffe, *Zhur. Obshchei Khim.* (J. Gen. Chem.) 21:1910-19 (1951); cf. C.A. 45, 3900b, 6281c. --Hydroxyfuchsones dyes with MeO groups ortho to hydroxyl or keto groups are acetylated normally forming corresponding acetoxytriphenylcarbinols, which are hydrolyzed by acids or bases to colored salts of the original hydroxyfuchsones dyes. The acetylation in the presence of HCl leads to 2 reactions simultaneously: one yields the above products while the second reaction leads to acid isomerization of the original dyes with subsequent acetylation of the probably formed hydroxyfluorene derivs. The concept of free radical dimers, as representative of the 2nd set of products (above), as postulated by Eidlitz and Postovskii (C.A. 43, 152c), is in error. 3,3'-Dimethoxybenzidine, AcO, and NaOAc refluxed 6 hrs. give 63% of an acetylation product, m. 136-8° (from EtOH), whose salts are colorless

in org. solvents, while mineral acids give violet color; hot aq. alkali gives blue color; boiling the Ac deriv. with Zn-AcOH yields a product that no longer gives color with acids or alkalis; analysis of the Ac deriv. indicates the compn. $C_{24}H_{20}O_4$. Acetylation with $AcCl-Ac_2O$ gives a small amt. of a substance, m. 239-32°, identical with the product of acid isomerization of 3,3'-dimethoxybenzidine, and the mother liquor yields the product, m. 136-11°, described in above expt. Acetylation of 3,3',3''-trimethoxybenzidine with $Ac_2O-NaOAc$ gives 60% product, m. 168°, corresponding to the normal triacetate; heated with acids or alkalis it gives violet color of rubrocol salts. Acetylation with $AcCl-Ac_2O$ gave the above product and a less sol. product, m. 208-8°, identical to the Postovskii-Eidlitz product, unchanged by acids; alkalis on heating yield a colorless soln.

Cl. M. Kozlovskii

CP

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Hydroxyfuchson dyes. XIV. Structure of the bisulfite derivatives of hydroxyfuchson dyes. I. S. Ioffe (Naval Med. Acad., Leningrad). *Zhur. Obshchei Khim. (Gen. Chem.)* 21, 1514-17(1951); cf. C.I. 45, 10504. — Alk. hydrolysis of acetyl derivs. of hydroxyfuchson dyes, which are the Ac derivs. of the carbinol derivs., results in formation of colorless salts of the carbinols. Similar hydrolysis of the bisulfite derivs. of hydroxyfuchson dyes leads directly to the colored salts of the dyes. Hence, the bisulfite derivs. are not the sulfite esters of the carbinols as proposed by Bayer (*Ber. 39, 57(1906)*) but are the result of reaction of the bisulfite with the carbonyl group of the dye and should be regarded as the α -hydroxysulfonic acids. Thus, the bisulfite deriv. of 3,3'-dimethoxybenzaurine in H_2O treated in the cold with 5% NaOH gives immediately a blue color, which disappears only slowly and reappears on warming. Similar treatment of the Ac deriv. gives murkiness, which goes over to a colorless soln., which acquires a weak blue color only after long standing, which is intensified by heating, but disappears on cooling. G. M. Kouslapoff

IOFFE, I. S.

"Hydroxyfuchsone dyestuffs. XV. Acetoxifuchsones." (p. 1677)

SO: Journal of General Chemistry (Zhurnal Obshchei Khimii) 1951, Vol 21, No 9.

IOFFE, I. S.

Ioffe, I. S., Bolen'kii, B. G.- "Investigation of hydroxyfuchsone dyes. XVI.
2-Hydroxyfuchsone." (p. 1437)

SO: Journal of General Chemistry, (Zhurnal Obshchei Khimii), 1952, Vol. 22, No. 8

IOFFE, I. S.

Catalysts

Chemical Abst.
Vol. 48 No. 3
Feb. 10, 1954
Organic Chemistry

Quinones. 1. Reaction of *p*-benzoquinone with *p*-acetamidobenzoic acid. *J. B. Long, Univ. Missouri, Chem. 23, 126-8 (1953).* A soln. of *p*-AcNHCO₂CH₃ in 500 ml. H₂O, was filtered, brought to the b.p., and treated gradually with 11 g. *p*-benzoquinone, causing a temporary orange-yellow color, with birth of colorless product; after 18 min. boiling the colorless soln. was filtered yielding 7.1 g. C₁₄H₁₁O₄N₂S, m. 182-3° (from AcOH or EtOH). The product is 4-acetamido-2',5'-dihydroxydiphenyl sulfone (I). Heated on water bath with Ac₂O and pyridine 1 hr. it gave 4-acetamido-2',5'-dipacetoxydiphenyl sulfone, 64%, m. 182-4° (from aq. EtOH); on soln. in alkalis and subsequent acylation the 2 Ac groups are lost and the original material, m. 283°, is regenerated. Similar reaction of *p*-methoxycarbonylaminobenzenesulfonic acid gave 67% 4-methoxycarbonylamino-2',5'-dihydroxydiphenyl sulfone (II), 210-11° (from H₂O); very stable to hot mineral acids, but hydrolyzed with hot alkalis. I refluxed with 10% HCl and evapd. gave 83% 4-amino-2',5'-dihydroxydiphenyl sulfone, m. 178-8° (from aq. EtOH); the same is obtained by hydrolysis of I in hot 10% NaOH. II similarly yields this substance by hydrolysis with 10% NaOH. The amino deriv. yields the *tri*-Ac deriv., m. 240-2°, when heated with Ac₂O/pyridine, while mere Ac₂O yields the *N*-Ac deriv. identical with above described. The amino deriv. forms HCl salt, m. 208-10°, which hydrolyzes rather readily in aq. soln. (C. M. Kosolapoff)

1-31-54

IOFFE, I. S.

Quinol. series. II. Chloromethoxyquinones. Ioffe and A. E. Sukhina. *Zhur. Obshch. Khim.* 23, 215-16 (1953); cf. C.A. 48, 1209c. Methoxylation of chloro-*p*-benzoquinone with MeOH-ZnCl₂ leads not only to introduction of MeO group, but also to displacement of the Cl by a 2nd MeO group. To 6 g. vanillin in 40 ml. 4% NaOH was rapidly added 40 ml. 6% H₂O₂, and the dark brown soln. treated after 1 hr. with 8 ml. 30% H₂SO₄ (8 ml.), chilled, then treated at -3° with 40 g. Na₂Cr₂O₇ in 75 ml. 20% H₂SO₄ over 3 hrs., followed by 2 hrs. at 0°, giving 3.0 g. (71%) methoxy-*p*-benzoquinone, m. 144° (from EtOH). Direct chlorination of vanillin in CHCl₃ gave 2,4,6-Cl₃(MeO)-C₆H₂CHO, m. 168°, which (6 g.) in 40 ml. 4% NaOH treated as described above with 40 ml. 6% H₂O₂ 30-40 min., acidified with 20% H₂SO₄, and chilled gave a ppt. of 6-chloro-2-methoxyhydroquinone, m. 140°, the mixt. can be directly oxidized with Na₂Cr₂O₇ at -5° as described, yielding 77% 6-chloro-2-methoxy-*p*-benzoquinone, orange, m. 168-6° (from EtOH). Similar treatment of 2,4,6-Cl₃(MeO)-C₆H₂CHO gave 60% 5-chloro-2-methoxy-*p*-benzoquinone, yellow, m. 172-3° (from EtOH), less sol. in AcOH and EtOH than the 6-Cl analog. The yield of 2,5-dimethoxy-*p*-benzoquinone by the Haeckel method (*Ber.* 34, 3334(1901)) is increased beyond 33% by addn. of oxidizing agents to the starting material in oxidation-reduction reactions, and the yields were even lower than without such addns. (FeCl₃, PbO₂). III. Chlorination of methoxyquinones. *Ibid.* 299-303. Passage of HCl into methoxy-*p*-benzoquinone (I) in CHCl₃ gave a blue-violet ppt., free of Cl, m. 238-40° (from PhNO₂), also formed on addn. of mineral acids to I in AcOH. The product has been described earlier (Erdman, C.A. 28, 1337). Thus I dimerizes in the presence of mineral acids. Passage of Cl through I in CHCl₃ yields a dichloride, which yellow at 100° and

(OVER)

decomp. 120°, losing HCl and yielding an orange melt, which resolidifies and m. 158-63°, being transformed to 6-chloro-2-methoxy-*p*-benzoquinone. Thus introduction of a MeO group reduces the stability of quinone dichlorides. *p*-Benzoquinone dichloride m. 146° without decomp., and only at 170-80° does it slowly lose HCl, yielding an unstable monochloroquinone (cf. Dismuth, *et al.*, C.A. 20, 1044). Chlorination of methoxy-*p*-benzoquinone must be done with pure Cl₂ for even traces of HCl lead to dimerization mentioned above. The heating of the methoxyquinone dichloride is best done by spreading the substance in a thin layer in a dish and heating to 120° in a thermostat. Although its decompn. can lead to 2 isomeric chloromethoxy-*p*-benzoquinones, the product actually obtained is the pure 6-chloro-2-methoxy isomer, m. 173°. The yield reaches 70%. The methoxyquinone dichloride is also unstable on heating in various solvents. Thus in hot aq. EtOH it loses HCl and yields up to 80% 6-chloro-2-methoxy-*p*-benzoquinone, m. 159°. Addn. of the dichloride to hot AcOH gave 6-chloro-2-methoxybenzoquinone, m. 172-3°, also formed in hot dry Me₂CO but in aq. AcOH and aq. Me₂CO the 6-Cl analog is formed. In hot H₂O the dichloride also yields the 6-Cl deriv. Heating the dichloride in aromatic hydrocarbons under dry conditions gives different results; the substance is not decompd. in boiling xylene for over 1 hr., while hot CCl₄ can be used as recrystn. solvent; the dichloride thus purified m. 120° (decompn.). A trace of H₂O immediately yields 6-chloro-2-methoxy-*p*-benzoquinone. G. M. E.

IOFFE, I. S.

"Investigation of quinones. Part 3. Chlorination of methoxy-quinone." Ioffe, I. S.,
Sukhina, A. F. (p. 299)

SO: Journal of General Chemistry (Zhurnal Obshchei Khimii) 1953, Volume No. 23, No.2.

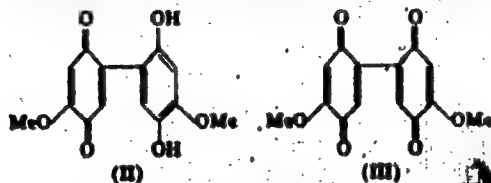
10FFE, 1-2

Chem Abs

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Organic Chem

✓ Quinone series. IV. Transformation of methoxyquinone under the influence of acids. I. S. 1976 and A. V. Sukhin. *Zhur. Obshchei Khim.* 23, 1370-6 (1953); *C. 1954*, 285; Riddiman, C.A. 28, 1337^o.—In contrast with other quinones, methoxyquinone (I), under the action of mineral acids, forms a condensation product with formation of a biphenyl link in a position para to the MeO group. Thus 10 g. I in 100 ml. warm AcOH, poured into 2 l. 1% HCl, gradually yields a blue ppt., which after 48 hrs. amounts to 95% II, m. 230° (crude), m. 236° (from pyridine), which is generally but sparingly sol. in org. solvents.



Reduction of 3 g. II with 3 g. Zn dust in refluxing AcOH gave, upon filtration, diln., and treatment with dil. NaHSO₄, a 60% yield of (5,5',4'(HO)MeOC₆H₄)₂, decomp. 210° (from dil. EtOH); *litro-Ac decomp.* from AcO-pyridine, m. 180-7° (from EtOH). II (3 g.) oxidized with 3 g. chromic acid in 300 ml. H₂O gave III, m. 212-14° (from AcOH). I (1 g.) in 10 ml. AcOH treated with 20 ml. concd. HCl gave after 48 hrs. 78% 5,5'-dihydroxy-4,4'-dimethoxy-3-chlorodiphenyl oxide, m. 220-1° (from AcOH), colorless, the diacetate (AcO-pyridine), m. 220-1° (from AcOH); also obtained in 65% yield from II in warm AcOH with concd. HCl, the blue color of II being discharged at 65-70°. III (6 g.) in 25 ml. AcOH refluxed with 25 ml. concd. HCl gave 86% 5,5'-dihydroxy-4,4'-dimethoxy-3,3'-dichlorodiphenyl oxide, m. 225-6° (from AcOH); diacetate, m. 225-3° (from AcOH).

O. M. Kozlov

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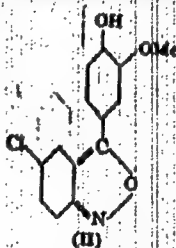
IOFFE, I. S.

CATALYST

Chemical Abst.
Vol. 48 No. 3
Feb. 10, 1954
Dyes and Textile Chemistry

(2) (Chem.)
Hydroxyfuchsons are series XVII. Condensation of isomeric nitrobenzaldehydes with gualacol. I. S. Ioffe and B. G. Belen'kii. *Zhur. Obshch. Khim.* 23, 1023-1024 (1953) *C. A.* 48, 2431k; 49, 8083g; 49, 8086c. $\text{m-O}_2\text{NC}_6\text{H}_4\text{CHO}$ (21 g.) and 38 g. gualacol, mtd. with dry HCl (3.5 g.) and kept in the cold 10 days in a closed flask, gave a mass of solid product, which was extr. several times with hot 2% Na_2CO_3 . The crude product treated with 60% EtOH and finally crystd. from 50% EtOH gave 3'-nitro-5,3'-dimethoxy-4,4'-dihydroxydiphenylmethane, m. 141.5°. The yield of crude product is 60.5% after the extr. with 60% EtOH ; yield of pure product is unstated. The crude product (5 g.), after the Na_2CO_3 treatment, was taken up in 50 ml. AmOAc and satd. with dry HCl , followed by a stream of N oxides (from $\text{NaNO}_2\text{-H}_2\text{SO}_4$) while maintaining the HCl stream. After 9 hrs. the mixt. was allowed to stand overnight, it deposited 68.8% crude 3'-nitro-5,3'-dimethoxy-4,4'-hydroxyfuchsono-HCl. With pure starting material the yield is 88%. The product is violet with green sheen and has no definite m.p. The HCl salt ground with excess 25% KOAc and 4 parts H_2O and allowed to stand overnight gave a ppt. of the hydrate form of the fuchsono, red solid. The latter (4.0 g.) heated with 60 ml. AcOH and cooled, gave 50.5% free fuchsono, red-orange, m. 230-9.5° (from AcOH); it is sol. in aq. acids and bases, giving red-violet soln. in acids and blue in bases. $\text{p-O}_2\text{NC}_6\text{H}_4\text{CHO}$ (4.9 g.) and 9.8 g. gualacol mtd. with dry HCl (0.7 g.) and kept in the cold 15 days, then treated with hot H_2O , gave crude yellowish green base, m. 10-60°, which (5 g.) was treated in AmOAc with HCl and N oxides as above, yielding 66% 3'-nitro-5,3'-dimethoxy-4,4'-hydroxyfuchsono-HCl, violet, without definite m.p. The latter (3.5 g.) with aq. KOAc gave the brick-red hydrate form of the fuchsono (3 g.), which crystd. from AcOH gave 48% free

fuchsin, red, m. 275-5.5°. sol. in acids with violet color, in bases with blue-green color. When 5 g. $\text{O}_2\text{NC}_6\text{H}_4\text{CHO}$ and 9 g. guaiacol were acid. with 1.1 g. dry HCl and kept 3 weeks there was formed a viscous orange mass. This kept overnight under 15 ml. 70% EtOH and dried gave 2.92 g. golden-yellow crystals, which could be crysdl. from various org. solvents; on recrystn. from AcOH it m. 181.5°. The product failed to undergo oxidation under the conditions used for the cases described above. Its compn. was $\text{C}_{17}\text{H}_{15}\text{ClN}_2\text{O}_4$ (I) indicating the condensation of 1 mol. of aldehyde with 1 mol. of guaiacol and simultaneous chlorination occurring. When the product was reduced with HCl-Zn dust and the filtrate was diazotized and coupled with 2-C₆H₄OH, a red dye was formed; with H-acid the dye was red-violet. The results indicate that I had the structure II, apparently formed by rearrangement of the intermediate $3,4\text{-(MeO)(HO)C}_6\text{H}_3\text{CH(OH)(C}_6\text{H}_4\text{NO}_2\text{)-}$.



G. M. Koenig

9-2-54
EJP

IOFFE, I.S.; SUKHINA, A.F.

Research in the field of quinones. Part 5. Chlorination of 2,5-dimethoxy-
quinone. Zhur.ob.khim. 23 no.10:1752-1757 0 '53. (MLRA 6:11)
(quinone)

100FF, 15.

Hydroxyfuchsin dyes. XVIII. Halogen derivatives of 3,3'-dimethoxybenzidine. I. S. Joffe and R. G. Pater. Zhur. Obshchei Khim. 23, 1635-1638 (1951); cf. C.A. 45, 1831. —Passage of dry HCl into 0.03 mole alcoholysis of 0.12 mole guaiacol, followed by storage of the mixture 1 week at room temp. in a closed vessel, treatment with hot H₂O and extraction with hot 10% aq. Na₂CO₃ gave 65-80% of the following leucobases, which were purified by soln. in EtOH and crystals from EtOH or AcOH: 3,3'-dimethoxy-4,4'-dihydroxy-1,1'-phenylene (halogen shown): 2'-chloro, decomp. 100-130°; 3'-chloro, decomp. 130-140°; 4'-chloro, m. 128-31°; 2'-bromo, decomp. 120-30°; 3'-bromo, decomp. 120-30°; 4'-bromo, m. 128-7°. These leucobases in AcOH were treated with dry HCl, then treated over 2 hrs. with a stream of N oxides containing NO₂, yielding a ppt. of the HCl salt of the dye in 20% yield: 3,3'-dimethoxy-4'-fuchsin (halogen shown): 2'-chloro, m. 164-5°; 3'-chloro, m. 147-5°; 4'-chloro, m. 202°; 2'-bromo, m. 171-2°; 3'-bromo, m. 163-4°; 4'-bromo, m. 202-4°. These salts are hydrolyzed by H₂O, yielding the hydrated forms of the hydroxyfuchsin; boiling these with AcOH, followed by distn. of the solvent in vacuo, gave the anhydrous forms of the dyes, which were purified by crystals from dry Cl₂ or AcOH; these were, resp.: 2'-chloro, orange-yellow, purple in acids, blue-violet in alkalis; 3'-chloro, orange-red, violet in acids, blue in alkalis; 4'-chloro, red, red-purple in acids, blue in alkalis; 2'-bromo, yellow-orange, violet in acids, blue-violet in alkalis; 3'-bromo, red, purple in acids, blue-violet in alkalis; 4'-bromo, red, purple in acids, blue in alkalis. G. M. H.

Ioffe, I. S.

USSR/Chemistry - Dyes

Card 1/1 Pub. 151 - 30/38

Authors : Ioffe, I. S., and Belen'kiy, B. G.

Title : Investigation of oxyfuchsona dyes. Part 19.- Effect of nitro-group and halogens on the indication properties of 3,3-dimethoxybenzaurin derivatives

Periodical : Zhur. ob. khim. 24/2, 343-352, Feb 1954

Abstract : The effect of nitro-groups and halogens in the benzene nucleus of 3,3-dimethoxybenzaurin on the indicating properties of dyes was investigated. It was found that the presence of these substitutes weakens the basic properties of dyes, their ability to form onic cations in strong acid media and reduces the stability of the cations. The presence of halogens in position relative to the central carbon atom, screens the latter and produces steric hindrances even during hydration of the dyes. The effects of the electron-acceptor and electron-donor nitro-groups on the conversion of 3,3-dimethoxybenzaurin derivatives into carbinol compounds are discussed. Four USSR references (1947-1953). Tables; graphs.

Institution : ...

Submitted : July 28, 1953

Ioffe, I. S.

USSR/Chemistry - Dyes

Card 1/1 Pub. 151 - 31/38

Authors : Ioffe, I. S., and Belen'kiy, B. G.

Title : Investigation of oxyfuchsons dyes. Part 20.- Effect of nitro-groups and halogens on the colority of 3,3'-dimethoxybenzaurin derivatives

Periodical : Zhur. ob. khim. 24/2, 353-361, Feb 1954

Abstract : The effect of substitutes on the colority of 3,3'-dimethoxybenzaurin derivatives containing nitro-groups and halogens in the benzene nucleus, was investigated. It was established that the presence of these substitutes has a definite effect of the colority of dyes in nonionized state, and in ionized state only in strongly acid and alkaline solutions. Nitro-groups and halogens cause a deepening of the color and onium cation of the dye in a strongly acid solution and a deepening of the dye anion in an alkaline solution. The conditions leading to maximum bathochromic effects of the nitro-groups and halogens are discussed. Ten references: 7-USSR; 1-USA and 1-English; 1-B. African (1900-1954). Tables; graphs.

Institution : ...

Submitted : July 28, 1953

Ioffe, I. S.

USSR/Chemistry - Condensation products

Card 1/1 : Pub. 151 - 26/37

Authors : Ioffe, I. S., and Khavin, Z. Ya.

Title : Investigation of quinones. Part 6.-Condensation of p-benzquinone with alpha-amino acids

Periodical : Zhur. ob. khim. 24/3, 521-527, Mar 1954

Abstract : Investigations showed that the reaction of p-benzquinone with alpha-amino acids is followed by oxidizing cleavage of the amino-acid and consequent formation of products of their condensation with o-benzquinone. The condensation products obtained from p-benzquinone with glycine and from p-benzquinone with alpha-alanine are listed. The physico-chemical properties of these condensation products are described. Ten references: 3-USA; 3-German; 1-Polish; 1-English; 1-Japanese and 1-French (1910-1950).

Institution :

Submitted : July 28, 1953

Ioffe, I. S.

USSR/Chemistry - Reaction processes

Card 1/1 : Pub. 151 - 27/37

Authors : Ioffe, I. S., and Khavin, Z. Ya.

Title : Investigation of quinones. Part 7.-Reaction of chlorine derivatives of p-benzoquinone with glycine esters

Periodical : Zhur. ob. khim. 24/3, 527-532, Mar 1954

Abstract : The reaction of chloroquinones with amines and with non-substituted quinone was investigated. The formation of a diamino-quinone derivative having both amino-groups in p-position relative to each other is described. The mechanisms and tendencies of such a chloroquinone reaction and its dependence upon the number of halogen atoms are discussed. Ten references: 5-German; 4-French and 1-USSR (1881-1954).

Institution :

Submitted : July 28, 1953

Ioffe I.S.

USSR.

V. Oul'povskii, VIII. Condensation of *p*-benzoquinone with
sulfanilamides. I. S. Ioffe, N. A. Filippova, and Z. Ya.
Kharin. *Izv. Akad. Nauk SSSR, Khim. Nauk* (1974), 4, 1365.
48: 13657. — Addn. of 2.2 g. *p*-benzoquinone to 100 ml. hot
H₂O to 2.5 g. sulfapyridine in 100 ml. EtOH followed by
refluxing 3 hrs. gave 2.08 g. 2,5-disulfapyridine-*p*-benzo-
quinone, m. above 300°. Similarly sulfathiazole gave 2,5-
disulfathiazole-*p*-benzoquinone, m. above 300°, while sulfa-
pyrimidine gave 2,5-disulfapyrimidine-*p*-benzoquinone. Car-
bazole gave 2,5-disulfacarbazole-*p*-benzoquinone, m. above 300°. *p*-Benzoquinone in 2 l. warm
H₂O was added to 3.6 l. cold H₂O and 0.5 l. aq. NaCl; when
the soln. reached room temp. it was treated with 9 g. sul-
fanilamide in 100 ml. EtOH and after 3 days at room temp.
gave 5 g. 2-sulfanilamide-*p*-benzoquinone, purified by extr.
with hot EtOH in which the bis-analog. was insol. while
dila. of the alc. ext. with H₂O gave the pure 2-sulfanilamide-*p*-
p-benzoquinone. Similarly was prepd. 2-sulfapyridine-*p*-
p-benzoquinone. Similarly was prepd. 2-sulfathiazole-*p*-
p-benzoquinone, sol. in EtOH and AcOH. In soln. both the
mono-derivs. are slowly transformed into insol. substances,
possibly polymerization or condensation products. IX.
Reaction of methoxyquinones with anilines. I. S. Ioffe and
S. R. Sgibina. *Ibid.* 705-9. — To hot soln. of 5 g. 2-meth-
oxy-*p*-benzoquinone (I) in 75 ml. EtOH was added 1.5 g.
PhNH₂ and the mixt. refluxed 2 hrs. and cooled, yielded 3.5
g. red 1-methoxy-5-anilino-*p*-benzoquinone, m. 100° (from
50% EtOH). This (1 g.) in 50 ml. hot AcOH treated with 1

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IOFFE, I. S.

1. PhNH_2 and refluxed 2 hrs. gave after hot filtration and cooling 1.2 g. red 2,5-dianilino-*p*-benzoquinone, m. above 300° (from PhNO_2); the same can be obtained in 1 step by using excess PhNH_2 . 1.5 g. in 75 ml. hot EtOH treated with 2.5 g. sulfanilamide and refluxed 2 hrs. gave 3 g. 2-methoxy-5-sulfanilamido-*p*-benzoquinone, red-brown, m. $270-81^\circ$ (from AcOH); on heating in AcOH it changed to 2,6-disulfanilamidobenzoquinone. Similarly 1 and sulfapyridine gave 2-methoxy-5-sulfapyridino-*p*-benzoquinone, red, decomp. 273° (from 85% AcOH); sulfathiazole, similarly gave red 2-methoxy-5-sulfathiazolo-*p*-benzoquinone, decomp. $245-7^\circ$ (from 60% AcOH). To 1 g. 2,6-dimethoxy-*p*-benzoquinone in 75 ml. hot AcOH was added 1 g. PhNH_2 and after 2 hrs. refluxing the mixt. gave 1 g. 2,6-dianilino-*p*-benzoquinone, m. above 300° . When 1 g. 2,6-dimethoxy-*p*-benzoquinone in 75 ml. hot AcOH was treated with 0.5 g. PhNH_2 and refluxed 2 hrs. there was formed after concn. and diln. with H_2O 0.3 g. 2-methoxy-5-anilino-*p*-benzoquinone, m. 160° .

G. M. Kozlovskii

IOFFE, I. S.

USSR/Chemistry

Card 1/1

Authors : Ioffe, I. S. ; and Sukhina, A. F.

Title : Investigation of quinones. Part 9.- Reaction of methoxyquinones with amines.

Periodical : Zhur. Ob. Khim. 24, Ed. 4, 705 - 709, April 1954

Abstract : The reaction of methoxyquinones with amines shows two trends:
1) addition of the amino radical to the non-substituted carbon atom of the quinoid nucleus and 2) displacement of the methoxyl group by the amino group. The amino radical subjected to the effect of the methoxyl group rapidly attaches itself to the non-substituted carbon atom provided the latter is in para-position relative to the amino group. Displacement of methoxyl group by still another amino radical at an aniline surplus is already much slower. Six references; 5 USSR since 1946; 1 English 1946; 2 German since 1891. Chemical formulas.

Institution :

Submitted : July 28, 1953

IOFFE, Isaac Solomonovich; KHAVIN, Z.Ya., redaktor; MERLIN, Ye.Ya., tekhnicheskiy redaktor.

[Organic chemistry] Organicheskaya khimiya. Iss-vo khim. lit-ry,
1956. 438 p. (MIRA 9:7)
(Chemistry, Organic)

5 (3)

AUTHORS:

Ioffe, I. S., Zal'manovich, M. Z.

SOV/79-29-8-51/81

TITLE:

N-Substituted Amides of Salicylic Acid and Its Derivatives.
I. Arylides of 3,5-Dichloro- and 5-Nitrosalicylic Acid

PERIODICAL:

Zhurnal obshchey khimii, 1959, Vol 29, Nr 8, pp 2682 - 2685
(USSR)

ABSTRACT:

Some arylides of salicylic acid are highly active disinfectants (Ref 1) (e.g. the "Anabial"). Particular attention is due to the synthesis of the parent compound of this group, the salicylanilide, by condensation of aniline with esters of the salicylic acid, e.g. with salol (Ref 8), where phenol is separated. This "salol method" is recommended as a general method of synthesizing various arylides of salicylic acid by heating salol with amines (Ref 10), in the medium of an inert solvent, e.g. trichlorobenzene. The experiments carried out by the authors, however, indicated that the solvent decreases the yield and only complicates the process. This reaction is shown to proceed quite smoothly when a mixture of salol and amine is fused together at 150-180° in an equimolar ratio, in which case at the beginning of the reaction the low-melting

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N-Substituted Amides of Salicylic Acid and Its Derivatives. I. Arylides of 3,5-Dichloro- and 5-Nitrosalicylic Acid SOV/79-29-8-51/81

salol acts as a solvent, and later on the resultant phenol. After termination of the reaction (1-2 hours) the phenol is removed by distillation, and the arylamide is purified from the alkaline solution by precipitation with acid and recrystallization from alcohol. In this way, yields of 80-95% were obtained in different arylides such as *m*-anisidide (Ref 8), *n*-phenetidide (Ref 9), *o*-nitroanilide and others (Refs 1,11,9). This method was also used for derivatives of salicylanilide with substituents in the salicyloyl nucleus. Different chlorine derivatives of the salicylanilide have so far been obtained by chlorination of this compound (Refs 12-14), while the degree of chlorination depends on the reaction conditions, and mixtures of different chlorinated chlorine derivatives are formed. The authors obtained easily the chlorine derivatives of salicylanilide in a pure state by fusing together the salol (and, accordingly, the chlorine salts) with aniline or chloro anilines. In this way, the 4'-chloro salicylanilide and 2',5'-dichloro salicylanilide (Ref 9) were obtained from

Card 2/3

N-Substituted Amides of Salicylic Acid and Its
Derivatives. I. Arylides of 3,5-Dichloro- and
5-Nitrosalicylic Acid

SOV/79-29-8-51/81

salol, and the 3,5,4'-trichloro salicylanilide and 3,5,2',5'-
tetrachloro salicylanilide (Ref 1) from 3,5-dichloro salol.
Tables 1 and 2 present further arylides of the 3,5-dichloro
salicylic acid and 5-nitrosalicylic acid synthesized in the
same way. There are 2 tables and 16 references, 2 of which
are Soviet.

ASSOCIATION: Voenno-meditsinskaya akademiya imeni S. M. Kirova (Military
Medical Academy imeni S. M. Kirov)

SUBMITTED: July 19, 1958

Card 3/3